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(71) Applicant: CANON KABUSHIKI KAISHA Ohta-ku, Tokyo (JP)

(72) Inventors:

 Kenmoku, Takashi Tokyo (JP)

- Sugawa, Etsuko Tokyo (JP)
- Tokyo (JP)
 Yano. Tetsuva
- Tokyo (JP)
- Mihara, Chieko Tokyo (JP)
- Imamura, Takeshi
- Tokyo (JP)
 Fukui, Tatsuki
 Tokyo (JP)
- (74) Representative: Weser, Wolfgang, Dr. Dipl.-Phys. Weser & Kollegen, Patentanwälte, Radeckestrasse 43
 - Radeckestrasse 43 81245 München (DE)
- (54) Polyhydroxyalkanoate having amide group and sulfonic group, method of producing the same, charge controlling agent containing the polyhydroxyalkanoate, toner binder, toner, and image formation method and image forming apparatus using the toner

(57) A polyhydroxyalkaneate comprises a unit of formula (1):-(O-CH((GH₂)_mSASO₂?\(\text{TO}\)_2C(-O))- wherein it is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and OC₂H₃; A represent a substituted or unsubstituted aliphatic hydrocarbon structure; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R, A and m in one unit can be different from them in another unit respectively. A method of producing the polyhydroxyalkanoate comprises the step of reacting a polyhydroxyalkanoate comprising a unit of formula [3]:-(O-CH((GH₂)_mB)CH₂C(G)), wherein m is an integer number selected from 1 to 8, and in the case where a plurality of units exist in the same molecule, m in one unit can be different from that in another unit, with at least one type of compounds of formula (19): PSA-1, SO_{A15} wherein Fi₁; is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH, and OC₂H₃ and A, is a substituted or unsubstituted aliphatic hydrocarbon structure, and in the case where a plurality of types of compounds axist in the same molecule, Fi₁; and A, in one unit can be different from them in another unit respectives.

Description

BACKGROUND OF THE INVENTION

5 Field of the Invention

[0001] The present invention relates to an innovative polyhydroxyalkanoate (may hereinafter be abbreviated as PHA) and a method of producing the same. Particularly, the present invention relates to a polyhydroxyalkanoate having hydrophilic proups and a method of producing the same.

[0002] In addition, the present invention relates to a charge controlling agent for use in electrophotography, electrostator recording, magnetic recording and the like, a toner binder, an electrostatic latent image developing toner, an image formation method using the toner, and an image forming appearatus for use therein. Particularly, the present invention relates to a charge controlling agent for use in electrophotography, electrostatic recording and electrostatic printing in a copying appearatus, a printer, a facesimile machine and the like for forming an image in such a manner that a toner image is previously formed on an electrostatic latent image carrier (hereinafter, referred to simply as image carrier) and thereafter the image is transferred onto an object transfer material, a toner binder, an electrostatic latent image developing toner, an image forming appearatus for use therefore.

Related Background Art

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10033 So far, many methods have been known for electrophotography, and those methods are generally carried out in such a manner that an electric latent image is formed on an image carrier (photoconductor) by a variety of means using a photoconductive substance, the latent image is then developed with a toner to form a visible image, and the toner image is transferred onto an object transfer material such as a paper as necessary, followed by fixing the toner image on the object transfer material by heat and/or pressure or the like to obtain a copy. For the method or visualizing the electric latent image, a cascade development method, a magnetic brush development method, a pressurizing development method and the like are known. Further, a method using a magnetic toner and a rotary development sleeve with a magnetic pole placed at the center thereof in which the magnetic toner is caused to fly from the development sleeve von the photoconductor by a magnetic field is also used.

20 [0004] Development systems for use in development of an electrostatic latent image include a two-component development system using a two-component type developer constituted by a toner and a carrier, and a one-component development system using a one-component type developer constituted only by a toner and using no carrier.

[0005] Here, the coloring fine particle generally celled as a toner has a binder resin and a coloring material as essential components, and in addition thereto, magnetic powders and the like as necessary. For the method for imparting an electric charge to the toner, the electrificability of the binder resin itself may be used without using a charge controlling agent, but in this method, charge stability with time and humidity resistance are compromised, thus making it impossible to obtain high quality images. Therefore, the charge controlling agent is usually added for the purpose of maintaining and controlline the charge of the toner.

[0006] Today, charge controlling agents known in the art include, for example, azo dye metal complexes, aromatic dicarboxylic-metal complexes and salicylic acid derivative-metal complexes for the negative friction charging agent. In addition, for the positive friction charging agent and addition, for the positive friction charging agent, nigrosine based dyes, triphenyimethane based dyes, various kinds of quaternary ammonium salts, and organic tin compounds such as dibutyl tin oxide are known, but toners containing these substances as the charge controlling agent do not necessarily fully satisfy quality characteristics required for the toner such as the electrificiality and stability with time depending on their compositions.

45 [0007] For example, a toner containing an azo dye metal complex known as a negative charge controlling agent has an acceptable charge level, but may have reduced dispersibility depending on the type of binder resin to be combined because the azo dye metal complex is a low-molecular crystal. In this case, the negative charge controlling agent is not uniformly distributed in the binder resin, the charge level distribution of the obtained toner is significantly lacking in sharpness, and the obtained image has a low gray-level, resulting in a poor image formation capability. In addition, the azo dye metal complex has a unique color tone, and is thus presently used only for toners having limited colors around black, and if the azo dye metal complex is used as a color toner, its lack in clarity as a coloring agent required for obtaining an image having a thigh level of resultement for the color tone is a serious problem.

[0008] In addition, examples of almost colorless negative charge controlling agents include aromatic dicarboxylicacid metal complexes, but they may be disadvantageous due to the fact that they are not perfectly colorless, and that they have low dispersibility peculiar to low-molecular-weight crystals.

[0009] On the other hand, nigrosine based dyes and triphenylmethane based dyes are presently used only for toners having limited colors around black because they are colored themselves, and may be poor in time stability of toners for continuous copying, in addition, conventional quaternary armonium salts may have insufficient humidity resistance

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when formed into toners, and in this case, the stability with time may be so poor that high quality images are not provided as they are repeatedly used.

[0010] In addition, in recent years, attention has been given worldwide to reduction of wastes and improvement of safety of wastes in terms of environmental protection. This problem applies to the field of electrophotography as well. That is, as imaging apparatuses have been widely used, the amounts of wastes of printed papers, discarded toners and copying papers have increased year by year, and the safety of such wastes is important from a viewpoint of protection of lobal environment. Pohytvdrovalkanate (PHA).

10011] Resins that can be decomposed with time by the action of microorganisms and the like, namely blodegradable resins are under development in terms of environmental protection, and for example, many types of microorganism have been reported to produce biodegradable resins having polyester structures (polyhydroxysikanoate: hereinafter abbreviated as PHA) and accumulate the resin in the cell (Non-patent Document 1). These polymers may be used for production of various kinds of products through melt processing as in the case of conventional plastics. In addition, these polymers have an advantage that owing to their biodegradability, they are fully decomposed by microorganism in the natural environment. And unlike many synthetic polymer corporunds, they never remain in the natural environment to cause contamination. In addition, they are also excellent in blocompatibility, and are expected to be applied as medical relaceble members and the like.

[0012] It is known that such PHA may various compositions and structures depending on the type of microorganism to be used for the production of the PHA, the culture medium composition and the culture condition, and higher of such have been conducted malnly on control of the composition and structure of PHA to be produced in terms of improvements of prometries of PHA.

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- [1] First, the biosynthesis of PHA obtained by polymerizing a monomer unit with a relatively simple structure such as 3-hydroxybutyric acid (hereinafter abbreviated as 3HB) is exemplified as follows:
 - (a) those containing 3HB and 3-hydroxyvaleric acid (hereinafter abbreviated as 3HV) (see Patent Documents
 - (b) those containing 3HB and 3-hydroxyhexanoic acid (hereinafter abbreviated as 3HHx) (see Patent Documents 5 and 6);
 - (c) those containing 3HB and 4-hydroxybutyric acid (hereinafter abbreviated as 4HB) (see Patent Document 7);
 - (d) those containing 3-hydroxyalkanoate having 6 to 12 carbon atoms (see Patent Document 8); and
 (e) biosynthesis using a single alighatic acid as a carbon source (the resulting product is almost same as those
 - (e) blosynthesis using a single aliphatic acid as a carbon source (the resulting product is almost same as those of (d)) (see Non-Patent Document 2).

They are all PHA composed of monomer units each having an alkyl group in the side chain, synthesized by β-oxidation of hydrocarbons and the like or synthesis of fatty acid from saccharides by microorganism, namely "usual PHA".

Such PHA has already found considerable applications with proven performance particularly in the field of agriculture, the biodegradable resin is used in muich files, horticulture materials, slow-releasable agricultural chemcolas, fertilizers and the like. Also, in the leisure industry, the biodegradable resin is used in fishing lines, fishing tackles, golf requites and the like.

[2] However, if considering a wide range of application as a plastic, it cannot be the above described that PHA is fully usable in terms of properties in the present situation. For further expanding the range of application of PHA, it is important to conduct a wide range of studies on the improvement of properties, and for this purpose, development and search of PHA including monomer units of a variety of structures is prerequisite. On the other hand, PHA with a substituent group introduced in the side chain ("unsusual PHA") can be expected to be developed as a "functional polymer" with very useful functions and properties originating from the introduced substituent group according to desired characteristics and the like. That is, it is also an important challenge to conduct of development and search of excellent PHA enabling such functionality and biodegradability to be compatible with each other. Examples of substituent groups include groups containing armatic rings (pheny) group, pheny group, etc.), unsaturated hydrocarbons, sets groups, glying orgues, oyang groups, halogenated hydrocarbons and epoxide. Among them, studies on PHA having an aromatic ring group conducted.

(a) PHA containing a phenyl group or its partially substituted group

- It is reported that Pseudomonas oleovorans produces PHA containing 3-hydroxy-5-phenylvaleric acid as a unit using 5-phenylvaleric acid as a substrate (see Non-Patent Documents 3 and 4).
- It is reported that Pseudomonas oleovorans produces PHA containing 3-hydroxy-5-(4'-tolyl) valeric acid

- as a unit using 5-(4'-tolyl) valeric acid as a substrate (see Non-Patent Document 5).
- It is reported that Pseudomonas oleovorans produces PHA containing 3-hydroxy-5-(2',4'-dinitrophenyi) valeric acid and 3-hydroxy-5-(4'-ntrophenyi) valeric acid as a unit using 5-(2',4'-dinitrophenyi) valeric acid as a substrate (see Non-Patent Document 6).

(b) PHA containing a phenoxy group or its partially substituted group

- It is reported that Pseudomonas oleovorans produces a PHA copolymer of 3-hydroxy-5-phenoxyvaleric acid and 3-hydroxy-9-phenoxynonanoic acid using 11-pheoxyundecanoic acid as a substrate (see Non-Patient Document 7).
- An invention relating to a homopolymer consisting of 3-hydroxy-5-(monofluorophenoxy) pentaneate (3H5 (MFP)P) units or 3-hydroxy-5-(difluorophenoxyi) pentanoate (3H5(MFP)P) units, a copolymer containing at least (3H5(MFP)P) units or (3H5(DFP)P) units; Pseudomonas putida synthesizing these polymers; and a method of producing the above described polymers using Pseudomonas species is disclosed, and it is described that as an advantage of the above invention, a long chain aliphatic acid having substituent groups can be metabolized to synthesize a polymer having a phenoxy group substituted with 1 or 2 fluorine atoms at the side chain terminal, and steroerogularity and water repellency are provided while maintaining a high meting point and good processability (see Patent Document 9).

Studies are conducted on cyano-substituents and nitro-substituents in addition to the fluorine-substituent described above.

It is reported that PHA containing 3-hydroxy-p-cyanophenoxyhexanoic acid or 3-hydroxy-p-nitrophenoxyhexanoic acid as a monomer until is produced with octanoic acid and p-cyanophenoxyhexanoic acid or p-nitrophenoxyhexanoic acid as substrates using a Pseudomonas olevorans ATCC 29347 strain and a Pseudomonas putida KT 2442 strain (see Non-Patent Documents 8 and 9).

These reports are useful in obtaining polymers each having an aromatic ring in the side chain of PHA and having properties derived therefrom unlike general PHA whose side chain contains an alkyl group.

[3] In addition, as a new category, studies are conducted for producing PHA having an appropriate functional group in the side chain and using the functional group to create a new function, beyond mere modification of properties of PHA.

- It is reported that PHA containing a unit having a vinyl group at the terminal of the side chain was synthesized, and was thereafter epoxidized, whereby PHA containing a highly reactive epoxy group at the side chain terminal could be synthesized (see Non-Patent Documents 10 and 11).
- In addition, it is reported that PHA containing a unit having a vinyl group at the terminal of the side chain was synthesized, and thereafter benzoyl peroxide was used with per-O-acetyl-1-thio-β-mattose, whereby PHA containing a sugar-chain could be synthesized, and that PHA containing a unit having a bromo group at the terminal of the side chain was synthesized, and thereafter diethyl amine was used with per-O-acetyl-1-thio-β-maltose, whereby PHA containing a usuar chain could be synthesized (see Non-Patent Document 12).

Application of Biodegradable Resin to Toner

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[0013] Application of a biodegradable resin to a binder resin particularly in production of toners is proposed in the field of electrophotography as well. For example, U.S. Patent No. 5,004,684 (Patent Document 10) discloses a toner having as its composition a biodegradable resin, particularly polyhydroxy burlor, acid and polyhydroxy valeric acid, a copolymer thereof or a blend thereof, in addition, Japanese Patent Application Laid-Open No. 6:289644 (Patent Document 11) discloses an electrophotographic toner particularly for heated roll fixation characterized in that all east the binder resin contains a plant based wax and a biodegradable resin (e.g. polyester produced by microorganism, and natural polymer material of plant or animal origin), and the above described plant based wax is added in the above described plant based wax is added in the above described plant for in an amount of 5 to 50% by weight.

[0014] In addition, Japanese Patent Application Laid-Open No. 7-120976 (Patent Document 12) discloses an electrophotographic toner characterized by containing a lactic add based rasis as a binder resin. In addition, Japanese Patent Application Laid-Open No. 9-274335 (Patent Document 13) discloses an electrostatic latent image developing coner.

characterized by containing a polyester resin obtained by dehydrating polycondensation of a composition containing lactic acid and tri- or higher functional oxycarboxylic acid and a coloring agent.

[0015] In addition, Japanese Patent Application Laid-Open No. 8-262796 (Patent Document 14) discloses an electrophotographic toner containing a binder resin and a coloning agent, characterized in that the binder resin is composed

of a biodegradable resin (e.g. aliphatic polyester resin), and the coloring agent is composed of non-water soluble pigments. In addition, Japanese Patent Application Laid-Open No. 9-281746 (Patent Document 15) discloses an electrostatic latent image developing toner characterized by containing an urethane-modified polyester resin obtained by cross-linking polylactic acid with a thi- or higher functional polyvalent isocyanate and a coloring agent.

- 5 [0016] Any one of the above described electrophotographic toners contains a biodegradable resin as binder resin, and is regarded to be effective to contribute to preservation of environments and the like.
 - [0017] However, reports of examples of using a biodegradable resin in the charge controlling agent have not been known, and there is a room for further improvement for contribution to preservation of environments.
- [0018] In addition to the above described documents, the content of Japanese Patent Application Laid-Open No. 2001-178484 (Patent Document 16) is herein incorporated.

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[Patent Document 1] Japanese Patent Publication No. 6-15604
[Patent Document 2] Japanese Patent Publication No. 7-14352
[Patent Document 3] Japanese Patent Publication No. 8-19227
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[Patent Document 4] Japanese Patent Application Laid-Open No. 5-7492

[Patent Document 5] Japanese Patent Application Laid-Open No. 5-93049
[Patent Document 6] Japanese Patent Application Laid-Open No. 7-265065

[Patent Document 7] Japanese Patent Application Laid-Open No. 9-191893 [Patent Document 8] Japanese Patent No. 2642937

20 [Patent Document 9] Japanese Patent No. 2989175

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[Patent Document 10] U.S. Patent No. 5,004,664

[Patent Document 11] Japanese Patent

Application Laid-Open No. 6-289644 [Patent Document 12] Japanese Patent Application Laid-Open No. 7-120975 [Patent Document 13] Japanese Patent

Application Laid-Open No. 9-274335 [Patent Document 14] Japanese Patent Application Laid-Open No. 8-262796 [Patent Document 15] Japanese Patent

Application Laid-Open No. 8-262/96 [Patent Document 15] Japanese Patent Application Laid-Open No. 9-281746 [Patent Document 16] Japanese Patent

Application Laid-Open No. 2001-178484 [Non-Patent Document 1] "Biodegradable Plastic

Handbook* Biodegradable Plastic Research Associate, N.T.S. Co., Ltd., p. 178-197 (1995)

[Non-Patent Document 2] Appl. Environ. Microbiol, 58 (2), p.746 (1992) [Non-Patent Document 3] Makromol. Chem., 191, p.1957-1965 (1990)

[Non-Patent Document 4] Macromolecules, 24, p.5256-5260 (1991)

[Non-Patent Document 5] Macromolecules, 29, p.1762-1766 (1996) [Non-Patent Document 6] Macromolecules, 32, p.2889-2895 (1999)

[Non-Patent Document 7] Macromol. Chem. Phys., 195, p.1665-1672 (1994)

[Non-Patent Document 8] Can. J. Microbiol., 41, p.32-43 (1995) [Non-Patent Document 9] Polymer International, 39, p.205-213 (1996)

[Non-Patent Document 9] Polymer International, 39, p.205-213 (1996) [Non-Patent Document 10] Macromolecules, 31, p.1480-1486 (1996)

ness and fragility. Therefore, PHA with improved melt processability has been desired.

[Non-Patent Document 11] Journal of Polymer Science: Part A: Polymer Chemistry, 36, p.2381-2387 (1998) [Non-Patent Document 12] Macromol. Rapid Commun. 20, p.91-94 (1999)

[0019] As described above, researches are being conducted for creating a new function, but only few successful cases have been reported. In particular, 3-hydroxybutylic acid has an advantage that it is completely decomposed by microorganism in the nature, but it has a problem in terms of melt processability because of its high crystallinity. Bard-

SUMMARY OF THE INVENTION

[0020] An object of the present invention is to provide an innovative polyhydroxyalkanoate with sulfionic group as a bydytophilic group or a derivative thereof introduced for improving melt processability, and a method of producing the same. In addition, the polyhydroxyalkanoate of the present invention is excellent in biocompatibility owing to its hydrophilic nature, and is expected to be applied as medicial flusible members and the like.

[0021] In addition, for solving the above described problems, the present invention provides a negatively end charge controlling agent being more contributable to preservation of environments and the like, and having high performance (high charge level, quick start of charge, excellent stability with time, and high environmental stability and improved dispersibility in the aspect of functionality, at one to inder containing the charge controlling agent, and all electrostatic latent image developing toner containing the charge controlling agent, and an image formation method and an image forming appearatus using the electrostatic leatent image developing toner. [0022] Thus, the inventors have devised the following invention as a result of vigorously conducting assertables for development of an innovative polyhydroxyalkanoate with a hydrophilic group introduced, which is believed to useful for improving melt processability. In addition, the inventors have vigorously conducted studies for developing a charge controlling agent having high performance and being substantially colorless, resulting in the achievement of the present invention.

[0023] That is, the outline of the present invention is as follows.

[1] A polyhydroxyalkanoate comprising a unit of formula (1):

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wherein R is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ ane OC₂H₅; A represents a substituted or unsubstituted alphatic hydrocarbon structure; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R, A and m in one unit can be different from them in another unit respectively.

[2] The polyhydroxyalkanoate according to item [1], comprising a unit of formula (2):

wherein R₁ is H or CH₃; R₂ is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and OC₂his, B, and C₁ each represent a substituted or unsubstituted aliphatic hydrocarbon structure; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R₁, R₂, B₁, C₁ and m in one unit can be different from them in another unit respectively.

[3] The polyhydroxyalkanoate according to item [2], comprising a unit of formula (3):

wherein R_3 is H or CH_3 : R_4 is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH $_3$ and CC_2 + R_4 : R_4 and R_5 are the represent a straight-chain or branched alkylene group having 1 to 8 carbon atoms; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R_3 , R_4 , R_5 , C_2 and m in one unit can be different from them in another unit respectively. (4) The polyhydroxyalknosted according to left mig. 3], comprising a unit of formula (4):

wherein R_5 is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH $_3$ and OC $_2$ H $_5$; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R_5 and m in one unit can be different from them in another unit respectively.

[5] The polyhydroxyalkanoate according to item [1], wherein m in formula (1) is an integer selected from the group consisting of 4, 6 and 8.

[6] The polyhydroxyalkanoate according to item [1], wherein m in formula (1) is an integer of 3 or 5.

[7] The polyhydroxyalkanoate according to item [1], comprising at least one of a 3-hydroxy-ω-alkanoic acid unit of formula (5):

wherein n is an integer number selected from 1 to 8; $R_{\rm G}$ contains a residue having a phenyl structure or a thienyl structure; and in the case where a plurality of units exist in the same molecule, n and $R_{\rm G}$ in one unit can be different from them in another unit resocitively and

a 3-hydroxy-ω-cyclohexylalkanolc acid unit of formula (6):

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wherein R_j is a substituent group in the cyclohexyl group selected from the group consisting of H_i CN_i NC_{ij} , as halogen atom, CH_{ij} CH_{ij} , CH_{ij} CH_{ij} , CH_{ij} CH_{ij} , CH_{ij} CH_{ij} C

[8] The polyhydroxyalkanoate according to item [7], wherein R₆ in formula (5) Is selected from the group consisting of the groups of the following formulas (7), (8), (9), (10), (11), (12), (13), (14), (15), (16) and (17):

unsubstituted or substituted phenyl groups of formula (7):

wherein R_{Ba} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, CH₂, C₂H₃, C₃H₃, CH=CH₂, COOR_{Ba}, (R_{Ba}, represents any one selected from the group consisting of H, Na and K), CF₃, CP₅, and C₃F₃, and in the case where a plurality of units exist in the same molecule, R_{8a} in one unit can be different from that in another unit;

unsaturated or saturated phenoxy groups of formula (8):

wherein R_g represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, CH₃, C₂H₅, C₃H₇, SCH₃, C₂H₅, C₃H₇, SCH₃, C₃H₅, C₃H₇, and in the case where a plurality of units exist in the same molecule, R_g in one unit can be different from that in another unit; unsubstituted or substituted benzoyl groups each of formula (9):

R₁₀ 0

wherein R_{10} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO_2 , CH_3 , C_2H_5 , C_3H_5 , C_3F_5 , C_2F_5 , and C_2F_7 , and in the case where a plurality of units exist in the same molecule, R_{10} in one unit can be different from that in another unit; unsubstituted or substituted phenylsulfanyl groups of formula (10):

wherein R_{114} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, ON, NO_2 , $COOR_{115}$, SO_2R_{116} , R_{116} represents any one selected from the group consisting of H, Na, K, CH_3 and C_3H_3 , and R_{116} represents any one selected from the group consisting of OH, ON_3 , ON_4 , a halogen atom, OCH_3 and OC_2H_3 , CH_3 , C_2H_3 , C_3H_3 ,

wherein R_{12a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO_a , $COOR_{12b}$, SO_a P_{12c} R_{12b} represents any one selected from the group consisting of H, Na, K, CH_3 and C_2H_3 , and R_{12c} represents any one selected from the group consisting of OH, ON_a , OK, a halogen atom, OCH_3 and OC_2H_3 , CH_3 , C_2H_5 , C_3H_7 , $(CH_3)_2$ -CH and $(CH_3)_3$ -C, and in the case where a plurality of units exist in the same molecule, P_{12a} in one unit can be different from that in another unit; P_{12a} P_{12a} in one unit can be different from that in another unit;

2-thienylsulfanyl group of formula (13):

2-thienylcarbonyl group of formula (14):

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unsubstituted or substituted phenylsulfinyl groups of formula (15):

wherein R_{13a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, COOR_{13b}(R_{13b}, represents any one selected from the group consisting of H, Na, K, CH₃ and C₂H₃), SO₂R_{13c}R_{10c}, represents any one selected from the group consisting of OH, Na, CK, a halogen atom, COH₃ and OC₂H₃), CH₃, C₂H₃, C₃H₇, (CH₃)₂·CH and (CH₃)₃·C, and in the case where a plurality of units exist in the same molecule, R_{13a} in one unit can be different from that in another unit; unsubstituted or substituted phenylsulfong reprose of formula (16):

wherein R_{14a} represents a substituent group in the aromatic ring selected from the group consisting of H_1 a halogen atom, CN_1 , NO_2 , $COOR_{14a}(H_{14a}$, represents any one selected from the group consisting of H_2 , N_1 , N_2 , N_3 , N_4 ,

, and in the case where a plurality of units exist in the same molecule, R_6 in one unit of formula (5) can be different from that in another unit.

[9] The polyhydroxyalkanoate according to item [1], wherein the number-average molecular weight is in the range of from 1000 to 1000000.

[10] A method of producing polyhydroxyalkanoate comprising a unit of formula (1), which comprises the step of:

reacting a polyhydroxyalkanoate comprising a unit of formula (18):

, wherein m is an integer number selected from 1 to 8, and in the case where a plurality of units exist in the same molecule, m in one unit can be different from that in another unit.

with at least one type of compounds of formula (19):

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wherein R_{15} is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH $_3$ and OC $_2$ H $_3$ and A $_1$ is a substituted or unsubstituted aliphatic hydrocarbon structure, and in the case where a plurality of types of compounds exist in the same molecule, R_{15} and A_1 in one unit can be different from them in another unit respectively.

[11] A method of producing polyhydroxyalkanoate comprising a unit of formula (2), which comprises the step of:

reacting a polyhydroxyalkanoate comprising a unit of formula (18) with at least one type of compounds of formula (20):

$$HS-B_3$$
 $N-C_3-SO_2R_{17}$

wherein R_{16} is H or CH_3 ; R_{17} is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH_3 and OC_2H_5 ; B_3 and C_3 are selected from substituted or unsubstituted aliphatic hydrocarbon structures; and in the case where a plurality of types of compounds exist in the same molecule, R_{16} , R_{17} , B_3 and C_3 in one unit can be different from them in another unit respectively.

- 40 [12] The method according to item [10], wherein the reacting step is carried out under basic condition.
 - [13] The method according to Item [12], wherein at least one selected from the group consisting of dimelyiamine, diethylamine, intrimethylamine, interthylamine, divulpamine, morpholine, piperidine, sodium hydroxide, sodium carbonate, potassium carbonate, sodium methylate and sodium ethylate is used as a basic actavite in the reacting step.
- 45 [14] A charge controlling agent controlling the charged state of powder, the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (1).
 - [15] The charge controlling agent controlling the charged state of powder according to item [14], the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (2).
 - [16] The charge controlling agent controlling the charged state of powder according to item [15], the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (3).
 - [17] The charge controlling agent controlling the charged state of powder according to item [16], the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (4).
 - [18] The charge controlling agent according to item [14], comprising at least one of a 3-hydroxy-ω-alkanoic acid unit of formula (5) and a 3-hydroxy-ω-cyclohexylalkanoic acid unit of formula (6).
- [19] The charge controlling agent according to item [18], wherein R₆ in chemical formula (5) is selected from the group consisting of formulae (7), (8), (9), (10), (11), (12), (13), (14), (15), (16) and (17).
 - [20] The charge controlling agent according to item [14], wherein the powder-is an electrostatic latent image de-

veloping toner.

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- [21] The charge controlling agent according to item [14], wherein the number-average molecular weight of the polyhydroxyalkanoate is in the range of from 1000 to 1000000.
- [22] A toner binder for use in an electrostatic latent image developing toner, the toner binder containing the charge controlling agent according to item [14].
 - [23] An electrostatic latent image developing toner containing a binder resin, a coloring agent and the charge
 - controlling agent according to item [14].

[24] A method for forming an image which comprises the steps of:

- applying a voltage to an electrification member from the outside to electrify an electrostatic latent image carrier. forming an electrostatic latent image on the electrified electrostatic latent image carrier, developing the electrostatic latent image by an electrostatic latent image developing toner to form a toner image on the electrostatic latent Image carrier, transferring the toner image on the electrostatic latent image carrier to a record material. and
- 15 fixing the toner image on the record material by heat, wherein an electrostatic latent image developing toner according to item [23] is used.
 - [25] The method according to item [24], wherein the transferring step comprises a first transferring step of transferring the toner image on the electrostatic latent image carrier to an intermediate transfer body and a second transferring step of transferring the toner image on the intermediate transfer body to the record material.
 - [26] An Image forming apparatus comprising a means for applying a voltage to an electrification member from the outside to electrify an electrostatic latent image carrier, a means for forming an electrostatic latent image on the electrified electrostatic latent image carrier, a means for developing the electrostatic latent image by an electrostatic latent image developing toner to form a toner image on the electrostatic latent image carrier, a means for trans-
- 05 ferring the toner image on the electrostatic latent image carrier to a record material, and a means for fixing the toner image on the record material by heat, wherein an electrostatic latent image developing toner according to item (23) is used.
- [27] The Image forming apparatus according to item [26], wherein the transferring means comprises a first transferring means for transferring the toner image on the electrostatic latent image carrier to an intermediate transfer 30 body and a second transferring means for transferring the toner image on the intermediate transfer body to the record material

BRIEF DESCRIPTION OF THE DRAWINGS

35 [0024]

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- FIG. 1 shows a ¹H-NMR spectrum chart of polyhydroxyalkanoate (PHA) produced in Example 1:
- FIG. 2 is a schematic explanatory view of an image forming apparatus used in Examples 23 to 40 and Comparative Examples 7 to 12:
- FIG. 3 is a sectional view of a principal part of a development apparatus for a two-component developer used in Examples 23 to 40 and Comparative Examples 7 to 12:
 - FIG. 4 is a schematic explanatory view having a reuse mechanism of a toner used in Examples 41 to 43 and Comparative Examples 13 to 15:
- FIG. 5 is a sectional view of a principal part of a development apparatus for a one-component developer used in 45 Examples 41 to 43 and Comparative Examples 13 to 15; FIG. 6 is an exploded perspective view of a principal part of a fixation apparatus used in the Example of the present
 - invention:
 - FIG. 7 is an enlarged sectional view of a principal part showing a film state of the fixation apparatus used in the Example of the present invention at the time when it is not driven; and
 - FIG. 8 is a schematic view showing a blow-off charge level measuring apparatus for measuring the charge level of the toner

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

[0025] The compound for use in the present invention has a basic skeleton as a biodegradability resin, and is therefore capable of being used for producing various kinds of products through melt-processing and the like as in the case of conventional plastics, and also has a remarkable characteristic such that it is decomposed by microorganism and involved in the material cycle in the natural world unlike synthetic polymers derived from oil.

[0026] The compound presented in the present invention, which is suitable as a charge controlling agent for use in an electrostatic latent image developing toner of the present invention will specifically be described. The compound for use in the present invention is a polyester resin having 3-hydroxyalkanoate as a monomer unit, namely polyhydroxyalkanoate having units each expressed by chemical formula (1).

[0027] Here, when such a compound is produced by a method comprising a step of producing the compound by microorganism, the compound presented in the present invention is an isotate polymer composed only of a R configuration, but is not particularly limited to the isotactic polymer and can be used for an atactic polymer as long as the object of the present invention can be achieved in terms of both properties and functions.

[0028] The polyhydroxyalkanoate of chemical formula (1) desired in the present invention is produced by a reaction between polyhydroxyalkanoate containing 3-hydroxy-e-bromoalkanole acid units each expressed by chemical formula (18), which is used as a startion material and at least one two of compound expressed by chemical formula (19).

[Method of producing polyhydroxyalkanoate containing units each expressed by chemical formula (18)]

[0029] The polyhydroxyalkanoate containing units each expressed by chemical formula (18) for use in the present invention can be produced using, but not limited to, a method of production by microorganism, a method of production by a gene-manipulated plant crop system and a method of production by chemical polymerization. Preferably, the method of production by microorganism is used.

[0030] A method of producing polyhydroxyalkanoate containing units each expressed by chemical formula (18) in the present invention in which microorganism is used will be described in detail.

[0031] If the production by microporghams in the present invention is used, any microorganism may be used as long as it is capable of producing a polyhydroxyalkanoate containing united each expressed by chemical formula (18) when cultured in a culture endium containing at least on type or formpound expressed by chemical formula (21), and one example thereof is a microorganism belonging to the Pseudomonas genus.

(In the formula, p is an integer number selected from 1 to 8.)

[0032] More particularly, the microorganism includes Pseudomonas cichoril YN2 (FERM BP-7379, Pseudomonas cichoril H46 (FERM BP-7374), Pseudomonas sessain P161 (FERM BP-7378) and Pseudomonas pudida P16 (FERM BP-7378). These four types of microorganisms are deposited at International Patent Organism Cepositary, National Institute of Advanced Industrial Science and Technology (former National Institute of Ediscience and Technology (former National Institute of Ediscience and Technology) and described in Japanese Patent Laid-Open No. 2001-178484 (Patent Document 18).

[0033] These microorganisms are capable of producing PHA containing a corresponding o-substituted-3-hydroxyalkanoic acid as a monomer unit using as a raw material a o-substituted-straight chain alkanoic acid substituted at the chain terminal with a six-membered ring atom group selected from the group consisting of a substituted or unsubstituted phenyl group, a substituted or unsubstituted phenoxy group and a substituted or unsubstituted cyclohexyl group, or a or-substituted-straight chain alkanoic acid substituted at the chain terminal with a five-membered ring atom group such as a thienyl group.

(Culture process)

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[0034] For usual culture of a microorganism for use in the method of producing PHA according to the present Invention, for example, production of preservation strains and propagation to secure the number of cells and the level of activity required for production of PHA, a culture medium containing components required to propagate the microorganism to be used is appropriately selected. For example, as long as the growth and survival of microorganism is not adversely affected, any type of culture medium such as a general natural medium (bouillon medium, yeast extract, etc.) and a synthetic medium with a nutrient source added therein may be used. Culture conditions including among other things temperature, ventilation and agitation conditions may be selected as appropriate depending on the type of microorganism to be used.

[0035] In the production method of the present invention, any culture medium may be used for the culture medium for use in the process of culturing a microorganism as long as the culture medium is an inorganic satt culture medium containing a phosphate and a nitrogen source such as an ammonium salt or nitrate, and in the process of producing

PHA in the microorganism, the productivity of PHA can be improved by adjusting the concentration of the nitrogen

[0036] In addition, nutrients such as an yeast extract, polypeptone and a meat extract can be added to the culture medium as a matrix for promoting the propagation of the microorganism. That is, peptides may be added as an energy source and a carbon source in the form of nutrients such as an yeast extract, polypeptone and a meat extract.

[0037] Alternatively, for the culture medium, saccharides, for example, aldoses such as glyceroladehyde, enythrose, arabinose, xylose, glucose, galactose, mannose and fructose, alditols such as glycerol, enythritol and xylitol, aldonic acids such as gluconic acid, uronic acids such as glucuronic acid and galacturonic acid, and disaccharides such as mallose, sucrose and lactose may be used as an energy source and a carbon source consumed with propagation of the microoranism.

[0038] Instead of the above described sechandes, organic acids or salts thereof, more specifically organic acids involved in the TCA cycle and organic acids derived from a biochemical reaction with less steps by one or two steps than the TCA cycle, or water soluble salts thereof may be used. As the organic acid or salt thereof, hydroxycarboxylic acids and oxocarboxylic acids such as pyruvic acid, oxalacetic acid, citric acid, isoclitric acid, ketoglutaric acid, succinic acid, multipact acid and lactic acid or water soluble salts thereof can be used. Alternatively, amino acids or salts thereof, for example amino acids such as asparaginic acid and glutamic acid or salts thereof and a group consisting of pyruvic acid, oxalacetic acid, citric acid, isocitric acid, ketoglutaric acid, succinic acid, fumaric acid, malic acid, lactic acid as lats thereof, and are added to the culture medium and dissolved therein. Alternatively, when the amino acid or salt thereof is added, it is more preferable that one or more types are selected from a group consisting of asparaginic acid, glutamic acid and salts thereof, and are added to the culture medium and dissolved therein. Alternatively, when the amino acid or salt thereof is added, it is more preferable that one or more types are selected from a group consisting of asparaginic acid, glutamic acid and salts thereof, and are added to the culture medium and dissolved therein. At this time, as reculter, all or part thereof can be added in the form of a water soluble salt to be dissolved uniformly without affection.

the pH of the culture medium.

[0039] It is desirable that the concentration of the above coexisting substrate added to the culture medium as a carbon source for growth of the microorganism and energy source for production of PHA is usually selected so that it is in the range of from 0.1 to 5% (ww), per culture medium. That is, for peptides, seat extracts, organic acids or salts thereof, amino acids or salts thereof, and saccharides that are used as the above coexisting substrates, one or more tyees thereof may be added, and at this time, it is destrable that the total concen-

tration of these added substrates is with in the above described range of total concentrations.

9 [0040] It is desirable that the content of the matrix for production of desired PHA, namely the compound expressed by general formula (21) is selected so that it is in the range of from 0.0005 to 1% (w/v), more preferably 0.001 to 0.2% (w/v) per cultural medium.

[0041] Any temperature at which microorganism strains to be used can suitably be propagated may be selected as a culture temperature, and an appropriate temperature is usually in the range of from about 15 to 37°C, more preferably from about 20 to 30°C.

[0042] Any culture method such as liquid culture and solid culture may be used for the culture as long as it allows propagation of micrographism and production of PHA. In addition, any type of culture method such as batch culture, ded-batch culture, semi-continuous culture and continuous culture may be used. Forms of liquid batch culture include a method of supplying oxygen while shaking the microorganism in a shake flask, and a method of supplying oxygen adopting a stirring ventilation sets with a lar frementer.

[0043]. For the method of making the microorganism produce and accumulate PHA, a two-step culture method in which the microorganism is cultured by two steps may be adopted other than the one-step culture method in which the microorganism is cultured in an inorganic salt culture medium containing a phosphate and a nitrogen source such as an ammonium sait or a nitrate with a matrix added therein in a predetermined concentration as described above, in this two-step culture method, the microorganism is once propagated sufficiently in the inorganic salt culture medium containing a phosphate and a nitrogen source such as an ammonium salt or a nitrate with a matrix added therein in a predetermined consentration as a primary culture, and thereafter cells obtained by the primary culture are relocated to a culture medium with a matrix added therein in a predetermined concentration after limiting the amount of nitrogen source such as ammonium childred contained in the culture medium, and are further cultured as a secondary culture, thereby making the microorganism produce and accumulate PHA. Use of this two-step culture method may improve the producitivity of desired PHA.

[0044] Generally, a produced PHA type polyester has reduced water solubility because of the presence of hydrophobic atom groups such as a bromaskly group derived from a 3-Hydroxy-e-bromasklaencia cald unit in the side bain, and is accumulated in cells of the microorganism capable of producing PHA, and therefore can easily be separated from the culture medium by collecting cells progragated by culture and involved in production and accumulation the desired PHA type polyester. After the collected cells are washed and dried, the desired PHA type polyester can be collected.

[0045] In addition, PHA is usually accumulated in cells of such a microorganism capable of producing PHA. For the

FP 1 336 635 Δ1

method of collecting desired PHA from these microorganism cells, a method that is usually used may be adopted. For example, extraction with organic solvents used nas chloratorm, dichloromethane and acetone is most convenient. Other than the above described solvents, dioxane, tetrahydrofuran and acetonitrile may be used. In addition, in a working environment in which use of any organic solvent is not preferred, a method in which in stead of solvent extraction, any one of a treatment by surfactants such as SDS, a treatment by expression as treatment by chemicals such as the superior of the surface of

[0046] As one example of inorganic salt culture media capable of being used in the production method of the present invention, the composition of the inorganic salt culture medium (M9 culture medium) used in Examples described later is shown below.

Composition of M9 culture medium:

[0047]

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Na₂HPO₄: 6.3 KH₂PO₄: 3.0 NH₄CI: 1.0 NaCI: 0.5

(by g/L, at pH=7.0).

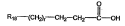
[0048] Further, for ensuring satisfactory propagation of cells and associated improvement of productivity of PHA, an essential trace element such an essential trace metal element should be added in an appropriate amount to an inorganic sati culture medium such as the above described M9 culture medium, and it is very effective to add about 0.3% (w/) trace component solution of which composition is shown below. The addition of such a trace component solution supclies a trace metal element for use in propagation of the microorganism.

30 Composition of trace component solution

[0049]

nitrilotriacetlo acid: 1.5; MgSO₄: 3.0; MnSO₄: 0.5; NaCl: 1.0; FeSO₂: 0.1; CaCl₂: 0.1; CoCl₄: 0.1; ZnSO₄: 0.1; CuSO₄: 0.1; AlK(SO₄)₂: 0.1; H₃BO₂: 0.1; Na₂MoO₄: 0.1; NiCl₅: 0.1 (g/L).

2 [0050] A matrix for producing desired PHA, namely at least one type of ω-substituted alkanotic acid compound expressed by chemical formula (22) or at least one type of ω-cytohexyliak knoic acid compound expressed by chemical formula (23) in addition to ω-bromoalkanotic acids each expressed by chemical formula (21) are made to coexist in the culture, whereby PHA containing 3-hydroxy-ω-substituted alkanotic acid units each expressed by chemical formula (5) or 3-hydroxy-ω-cytohexyliakanotic acid units from chemical formula (21) can be produced. The contents of ω-bromoalkanotic acid expressed by chemical formula (21), os-usbstituted alkanotic acid compound expressed by chemical formula (21), os-usbstituted alkanotic acid compound expressed by chemical formula (23) are selected so that they are in the range of from 0.0005 to 1% (w/v), more preferably from 0.001 to 2.5% (w/v) per culture medium, respectively.



r=1-8 (22)

(r is an integer number selected from 1 to 8; R₁₈ includes a residue having any one of a phenyl structure and a thieriest structure, and represents any one selected from the group consisting of florabulas (7, (8), (9, (10), (11), (12), (13), (14), (15), (16) and (17), and in the case where a plurality of types of compounds exists, R₁₈ and r in one unit can be different from them in another respectively.

$$R_{19}$$
 (CH₂)s - CH₂ - CH₂ OH (23)

(In the formula, R₁₉ represents a substituent group in the cyclohexyl group, and R₁₉ is selected from the group consisting of H, CN, NO₂, a halogen atom, CH₃, C₂H₅, C₃H₇, CF₃, C₂F₅ and C₃F₇, s is an integer number selected from 0 to 8.

(Method of producing polyhydroxyalkanoate expressed by chemical formula (1))

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- [0051] The reaction between the polyhydroxyalkanoate having units each expressed by the chemical formula (18) in the present invention and the compound expressed by chemical formula (19) will be described in detail.
- [0052] The compound expressed by chemical formula (19) for use in the present invention is used in the amount in the range of from 0.1 to 50.0 times in mole, more preferably from 1.0 to 30.0 times in mole as large as the amount of unit expressed by chemical formula (18) to be used as a starting material.
- [0053] The reaction in the present invention is preferably made to proceed under a basic condition. For the base, amines such as druthly aimine, diethyl amine, intentibyl amine, divuly aimine, morpholine and piperidine, alkali hydroxide metals such as sodium hydrate and potassium hydrate, alkali carbonate metals such as sodium carbonate, alkali and sodium carbonate, alkali and sodium carbonate, alkali metal alcoholates such as sodium methylate and sodium ethylate, sodium hydride and the like may be used. Particularly, diethyl amine, and dibutyl amine are preferably used. The amount of base to be used is in the range of from 0.1 to 100.0 times in mole, more preferably from 0.5 to 50.0 times in mole as large as the amount of unit expressed by chemical formula (18).
- [0054] A solvent may be used as required in the reaction of the present invention. Solvents to be used include hydrocarbons such as hexane, cyclohexane and heptane, ketones such as acetone, methyl eithyl ketone, ethers such as dimethyl ether, diethyl ether, tetrahydrofuran, halogenated hydrocarbons such as dichloromethane, chloroform, carbon tetrachloride, dichloroethane, and tichloroethane, aromatic hydrocarbons such as benzene and toluene, and aprotic polar solvents such as Ni-dimethyl formarindide particularly sulfoxide. Ni, V-dimethyl formarindide particularly preferably used. The amount of solvent to be used may be determined as appropriate according to the type of starting material, the type of base, the reaction condition and the like.
- [0055] In the method of the present invention, the reaction temperature is not particularly limited, but is usually in the range of from 0°C to the boiling point of the solvent. In the case where the reaction temperature is higher than a room temperature, however, the ester linkage of the main chain may be cut, and it is therefore more preferable that the reaction is made to proceed at a temperature of about 0 to 30°C.
 - [0056] In the method of the present invention, the reaction time cannot be determined unconditionally, but is usually in the range of from 1 to 48 hours.
- 45 [0057] In the present invention, the reaction solution containing the polyhydroxyalkanoate of chemical formula (1) produced in this way can be removed using distillation as a normal method. Alternatively, the reaction solution can be collected by mixing the reaction solution uniformly with a solvent insoluble in the polyhydroxyalkanoate expressed by chemical formula (1) to represent the desired ophydroxyalkanoate expressed by chemical formula (1) using a solvent such as water, an alcohol such as methanol and shanol, and an either such as dimethyl either, diethyl either diethyl either and tetrahydrofuran. The polyhydroxyalkanoate of chemical formula (1) obtained in this way may be isolated and purifiled if necessary. The isolation and purification method is not particularly limited, and a method in which a solvent insoluble in the polyhydroxyalkanoate expressed by chemical formula (1) is used to reprecipitate the polyhydroxyalkanoate, a method using oliving may be used.
 - [0058] Furthermore, in the reaction in the present invention, the reaction solvent, the reaction temperature, the reaction time and the purification method should not be limited to those described above.
 - [0059] In addition, the present invention is a charge controlling agent containing the above described polyhydroxyalkanoate, and turther an electrostatic latent image developing toner containing the charge controlling agent. The present invention is turther an image formation method comprising an charging step of applying a voltage to a charge

member from the outside to uniformly charge an electrostatic latent image carrier for the above electrostatic latent image developing lone, a development sisp of forming a toner image on the electrostatic latent image carrier, a transfer step of transferring the toner image on the electrostatic latent finage carrier to a transfer material via or not via an intermediate transfer body, and a heat-fixation step of fixing by heat the toner image on the transfer material, and also an image forming apparatus comprising means corresponding to respective steps of the method, namely charging means, development means, transfer means and heat-fixation means.

[0060] The polyhydroxyalkanoate for use in the present invention has good compatibility with the binder resin and excellent compatibility particularly with polysester type binder resin. Since the toner containing the polyhydroxyalkanoate of the present invention has a high specific charge level and is excellent in stability with time, it provides clear images with stability in image formation with electrostatic recording even after being stored for a long time period, and the toner can be produced for both negatively charged black toners and color toners because of its colorlessness and negativeetertrifiability.

[0061] In addition, by properly selecting the type and composition ratio of monomer units constituting the polyhydroxyalkanous of the present invention, wide range compatibility control is made possible. If a realism composition in which the charge controlling agent is put in micro-phase separation state in a toner binder, no electric continuity is formed in the toner so that electric charge can stably be maintained. In addition, the polyhydroxyalkanoate of the present invention contains no heavy matela, and therefore when the toner is produced by suspension polymerization or emulsion polymerization, polymerization inhibition caused due to the presence of heavy metals, as found in the case of a metal-containing charge controlling agent, does not occur, thus making it possible to produce a toner with stability.

(Addition of PHA to toner)

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[0082] In the present invention, the method for adding the above compound to a toner may be a method of internal addition to the toner. The addition amount of the internal addition is generally 0.1 to 50% by weight, preferably 0.3 to 30% by weight, and turther preferably 0.5 to 20% by weight as the weight ratio of the toner binder and the charge controlling agent. If it is lower than 0.1% by weight, the improvement degree of the charging property of the toner is insignificant and thus not preferable. Whereas, if it is higher than 50% by weight, it is not preferably from an economical point of view. Further, in the case of the external addition, the weight ratio of the toner binder and the charge controlling agent is preferably 0.01 to 5% by weight, and it is particularly preferable that the compound is mechanochmically fixed on the surface of the toner. In addition, the compound presented in the present invention may be used in combination of a known charge controlling agent.

[0033] The number-average molecular weight of the above-described compound of the present invention is usually 1000 to 1000000, preferably 1000 to 500000. If it is less than 1000, the compound is completely compatible with the toner binder to make it difficult to form a discontinuous domain, resulting in an insufficient charge level, and the fluidity of the toner is adversely affected. Further, if it is higher than 1000000, dispersion in the toner becomes difficult.

[0064] The molecular weight of PHA of the present invention can be measured as a relative or absolute molecular weight. It can conveniently be measured by, for example, SPC (ge) permeation orbimoteroptary). For the specific measurement process of GPC, PHA is dissolved in a solvent capable of dissolving the PHA in advance, and a measurement is made with a similar mobile phase. For the detector, a differential refraction detector (II), an uttraviolat detector (IV) or the like may be used depending on the PHA to be measured. The molecular weight is determined as a relative comparison with a standard sample (polystyrene, polymethylmethacylate, atc.). The solvent may be selected from solvents capable of dissolving a polymer such as dimethylmemamide (DMF), dimethylsulfoxide (DMSO), chloroform, tetrahydrofuran (THF), toluene, hexafluoroisopropanol (HFIP). In the case of a polar solvent, a measurement can be made by addition of salt. In addition, in the present invention, compounds presented in the present invention with the ratio (Mw/Mn) of the weight average molecular weight (Mw) to the number-average molecular weight (Mn) measured as described above being in the range of from 1 to 10 are preferably used.

[0065] In the present invention, the compound presented in the present invention has a melting point preferably in the range of from 20 to 150°C, especially preferably from 40 to 150°C, or so no melting point but a glass transition temperature in the range of from 20 to 150°C, especially preferably from 40 to 150°C. If the foregoing melting point is lower than 20°C or the glass transition temperature with no melting point is lower than 20°C, the fluidity and the storage property of the toner are often adversely affected. Whereas if the foregoing melting point is higher than 150°C, the charge controlling agent becomes difficult to be kneaded with the toner and the charge level distribution becomes broad in many cases.

[0066] To measure the moiting point Tm and the glass transition temperature Tg in this case, a high precision and internally heating input compensation type differential scanning calorimeter, for example, DSC-7 manufactured by Perkin Elmer Co, may be employed.

[0067] Regarding the toner binder and the electrostatic latent image developing toner of the present invention, the weight ratio of the toner binder and the charge controlling agent is generally 0.1 to 50% by weight, preferably 0.3 to

FP 1 336 635 Δ1

30% by weight, and more preferably 0.5 to 20% by weight. Regarding the composition ratio of the electrostatic latent image developing toner of the present invention, generally the foregoing charge controlling agent is in the range of from 0.1 to 50% by weight, the toner binder is in the range of from 20 to 95% by weight, and a coloring material is in the range of from 0.1 to 50% by weight, and a coloring material is in the range of from 0.1 to 50% by weight with respect to the weight of the toner and based on the necessity, a magnetic powder (a powder of a ferromagnetic metal such as iron, cobat, nickel and the like and a compound such as magnetite, hematite, ferrite and the like) functioning as a coloring material may be added in an amount not more than 60% by weight. Further, various additives (a lubricant (polyterafluoroethylene, a bwer motecuts weight) polyting in, an alphatic acid or its metal saft or amide, and the like) and other charge controlling agents (metal-containing azo dye, metal salytet, etc.) may be contained. In addition, in order to improve the fluidity of the loner, a phydrophotic colloidal silies fine powder may also be employed. The amounts of these additives are generally not more than 10% by weight on the bases of the toner weight.

[0088] In the toner of the present invention, it is preferable for at least some of the toner binder to form a continuous phase and at least some of the charge controlling agent to form discontinuous domain. As compared with the case where the charge controlling agent to favor discontinuous domain. As compared with the case where the charge controlling agent than a complete compatibility with the toner binder without forming the discontinuous domain, the added charge controlling agent is easily exposed to the surface and effective even in a small amount. The dispersion particle diameter of the domain is preferably 0.01 to 4 µm and more preferably 0.05 to 2 µm. If it is bigger than 4 µm, the dispersibility becomes insufficient and the charge level distribution becomes broad and the transparency of the toner is deteriorated. Whereas, if the dispersion particle dismeter is smaller than 0.01 µm, it becomes similar to the case where the charge controlling agent has complete compatibility with the binder without forming discontinuous domain, a large amount of the charge controlling agent is required to be added. That at least some of the foregoing charge controlling agent forms the discontinuous domain and the dispersion particle size can be observed by observing a specimen of the toner with a transmission electron microscope. In order clearly observe the interface, it is also effective to carry out observation of a toner specimen by electron microscope after the specimen is dyed with ruthenium tetraxide, a multimust retraxide, and the like.

[0069] Further, for the purpose of reducing the particle diameter of the discontinuous domain formed by the compound presented in the present invention, a polymer compatible with the compound presented in the present invention and also with the toner binder may be added as a compatible agent. The compatibility enhancing agent is, among other things, a polymer comprising mutually graft- or block-polymerized polymer chains containing at least 50% by mol of monomers having practically similar structure to that of the constituent monomers of the compound presented in the present invention and polymer chains containing at least 50% by mol of monomers having practically similar structure to that of the toner binder. The amount of the compatible agent to be used is generally not more than 30% by weight and preferably 1 to 10% by weight, with respect to the compound presented in the present invention.

(Other constituent materials)

[0070] Other constituent materials constituting the electrostatic latent image developing toner of the present invention will be described below.

(Binder resin)

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[0071] At first, any resin may be used as the binder resin without any particular restrictions if it is generally used for production of a toner. Also, the charge controlling agent of the present invention may previously be mixed with the binder resin to be used as a toner binder composition of the present invention having charge controlling capability before production of the toner. For example, styrene-based polymers, polyester-based polymers, polyester-based polymers, polyester-based polymers, polyester-based polymers, and they are used alone or while being mixed with one another.

[0072] The styrene-based polymers may be styrene-(meth)acrylic acid ester copolymers and copolymers of these copolymers with other monomers copolymerizable with them; copolymers of styrene with diene type monomers (but-diene, isoprene and the like) and copolymers of these copolymers with other monomers copolymerizable with them; and the like. The polyester-based polymers may be condensation polymerizable products of aromatic dientoxylic acid and aromatic diel alikylene oxide addition products and the like. The epoxy-based polymers may be reaction products of aromatic diels and epichlorohydrin and their modified products. The polyelen-based polymers may be reaction products of aromatic diels and epichlorohydrin and their modified products. The polyelen-based polymers may be addition products of aromatic diels with them. The polyurethane-based polymers may be addition polymerization products of aromatic diisocyanates and aromatic diel atkiviene oxide addition products and the like.

[0073] Practical examples of the binder resin to be employed in the present invention are polymers of the following polymerizable monomers or their mixtures, or copolymerization products produced from two or more kinds of the following polymerizable monomers. Such polymers are more particularly, for example, styrene-based polymers such as

styrene-acrylic acid copolymer, styrene-methacrylic acid copolymer and the like; polyester-based polymers; epoxybased colymers; polyolefin-based polymers; and polyurethane-based polymers and they are preferably used.

[0074] Practical examples of the polymerizable monomers are styrene and its derivatives such as styrene, o-methvistyrene, m-methylstyrene, p-methylstyrene, p-methoxystyrene, p-phenylstyrene, p-chlorostyrene, 3,4-dichlorostyrene, p-ethylstyrene, 2,4-dimethylstyrene, p-n-butylstyrene, p-tert-butylstyrene, p-n-hexylstyrene, p-n-octylstyrene, pn-nonylstyrene, p-n-decylstyrene, p-n-dodecylstyrene, and the like; ethylenic unsaturated monoolefins such as ethylene, propylene, butylene, isobutylene and the like; unsaturated polyenes such as butadiene and the like; vinyl halides such as vinyl chloride, vinylidene chloride, vinyl bromide, vinyl fluoride and the like; vinyl esters such as vinyl acetate, vinyl propionate, vinyl benzoate and the like; α-methylene aliphatic monocarboxylic acid esters such as methyl methacrylate, ethyl methacrylate, propyl methacrylate, n-butyl methacrylate, isobutyl methacrylate, n-octyl methacrylate, dodecyl methacrylate, 2-ethylhexyl methacrylate, stearyl methacrylate, phenyl methacrylate, dimethylaminoethyl methacrylate, diethylaminoethyl methacrylate, and the like; acrylic acid esters such as methyl acrylate, ethyl acrylate, nbutyl acrylate, isobutyl acrylate, propyl acrylate, n-octyl acrylate, dodecyl acrylate, 2-ethylhexyl acrylate, stearyl acrviate, 2-chloroethyl acrylate, phenyl acrylate, and the like; vinyl ethers such as vinyl methyl ether, vinyl ether, vinyl isobutyl ether, and the like; vinyl ketones such as vinyl methyl ketone, vinyl hexyl ketone, vinyl isopropenyl ketone, and the like; N-vinyl compounds such as N-vinylpyrrole, N-vinylcarbazole, N-vinylindole, N-vinylpyrrolidone and the like; vinyl naphthalenes; acrylic acid or methacrylic acid derivatives such as acrylonitrile, methacrylonitrile, acrylamide and the like; esters of the above-described α, β-unsaturated acid; diesters of bibasic acid; dicarboxylic acids such as maleic acid, methyl maleate, butyl maleate, dimethyl maleate, phthalic acid, succinic acid, terephthalic acid, and the like; polyols compounds such as ethylene glycol, diethylene glycol, triethylene glycol, 1,2-propylene glycol, 1,3-propylene glycol, 1,4-butanediol, 1,6-hexanediol, bisphenol A, hydrogenated bisphenol A, polyoxyethylene-modified bisphenol A and the like: Isocyanates such as p-phenylene diisocyanate, p-xylylene diisocyanate, 1,4-tetramethylene diisocyanate, and the like; amines such as ethylamine, butylamine, ethylenediamine, 1,4-diaminobenzene, 1,4-diaminobutane, monoethanolamine, and the like; epoxy compounds such as diglycidyl ether, ethylene glycol diglycidyl ether, bisphenol A glycldyl ether, hydroguinone glycldyl ether, and the like.

(Cross-linking agent)

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[0075] In the case of producing the binder resin to be employed in the present invention, based on the necessity, the following cross-inking agent may be used. Examples of a bifunctional cross-inking agent are diviny/benzene, bis (4-eory)dxypolyethoxypheny/lpropane, ethylene glycol diacrylate, 1,3-butylene glycol diacrylate, 1,4-butanediol diacrylate, 1,3-bexandiol diacrylate, persently glycol diacrylate, diethylene glycol diacrylate, triethylene glycol diacrylate, because glycol diacrylate, triethylene glycol diacrylate, because glycol diacrylate, because glycol diacrylate, because glycol diacrylate, because glycol diacrylate, polypropylene glycol dia

[0076] Examples of bi- or higher polytunctional cross-linking agent are pentiaeryhritot triacrylate, trimethylolethane triacrylate, triallyl accyanurate, triallyl accyanurate, triallyl trimellate, dairy chlorendate, and the like.

Polymerization Initiator

1,0077 In the case of producing the binder resin to be employed for the present invention, the following polymarization intillators may be used based on the necessity; for example, tert-buty peroxy2-ethylhexanoate, cumine perpivalate, tert-buty peroxydiautate, benzey) peroxide, tert-buty peroxydiautate, benzey) peroxide, tert-buty peroxydiautate, benzey) peroxide, zer-azobis (-demony-azobis (-de

(Other biodegradable plastics)

[0078] In addition, in the present invention, biodegradable plastics are preferably used. Examples of the biodegradable plastics are "Ecostar", "Ecostar plus" (produced by Hagwara Industries, Inc.), "Biopole" (produced by Monant Co.), "Ajicoat" (Ajinomoto Co., Ltd.), "Placcof", "Polycaprolectore" (produced by Daicel Chem., Ind., Ltd.), "Bionoile" (produced by Showa Highpolymer Co., Ltd.), "Lacty" (produced by Shimadzu Corporation), "Lacea" (produced by Misui Chemicals Inc.) and the Company of the

[0079] It is preferable for the combinations of the binder resin and the charge controlling agent of the present invention that the structure of the polymer chain of the charge controlling agent are similar to each other as much as possible. If the structure of the polymer chain of the charge controlling agent are similar to each other as much as possible. If the structure of the polymer of the binder resin and the polymer structure of the polymer chain of the charge controlling agent are considerably dissimilar to each other, the charge controlling agent are similar to each other, the charge controlling agent are structured to the polymer chain of the charge controlling agent are such as the charge controlling agent are considerable as the charge controlling agent are consi

[0080] The weight ratio of the charge controlling agent of the present invention to be internally added to the binder resin is generally 0.1 to 50% by weight, prefereably 0.3 to 30% by weight, and more prefereably 0.5 to 20% by weight, the the weight ratio of the charge controlling agent to be internally added is lower than 0.1% by weight, the charge level becomes low and if the weight ratio is higher than 50% by weight, the charge stability of the toner is deteriorated.

(Coloring agent)

- (20 [0081] Any coloring agent generally used for production of a toner may be used as the coloring agent composing the electrostatic latent image developing toner of the present invention without particular restrictions. For example, carbon black, titanium white, and any other pigment and/or dye may be used. For example, in the case the electrostatic latent image developing toner of the present invention is used for a magnetic color toner, examples of the coloring agent to be employed are C.I. Direct Red 1, C.I. Direct Red 4, C.I. Acid Red 1, C.I. Basic Red 1, C.I. Mordant Red 30, 192 (J. Direct Red 1). Direct Red 1, C.I. Direct Red 1, C.I. Basic Red 1, C.I. Basic Red 1, C.I. Mordant Red 30, 193 (J. Direct Red 1). Direct Green 6, C.I. Basic Red 1, C.I. Ba
- [0082] In the case the electrostatic latent image developing toner of the present invention is used for a two-component type full color toner, the following coloring agents can be used. For example, coloring pigments for magenia toners are 2C. I. Pigment Red 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 21, 22, 23, 30, 31, 32, 37, 38, 39, 40, 41, 48, 49, 50, 51, 52, 53, 54, 55, 57, 56, 60, 63, 64, 68, 81, 83, 87, 88, 89, 90, 112, 114, 122, 123, 163, 202, 206, 207, 209, CL. Pigment Violet 19, CJ. Vat Red 1, 2, 10, 13, 15, 23, 29, 35 and the little of the color of
- S5, 38, 37, 38, 39, 40, C.I. Basic Violet 1, 3, 7, 10, 14, 15, 21, 25, 26, 27, 28, and the like.
 [0884] As other coloring pipments examples of oyen coloring pipments ex. C.I. Pigment Blue 2, 3, 15, 16, 17, C.I.
 Val Blue 6, C.I. Acid Blue 45, copper-phthalocyanine pigments having a phthalocyanine skeleton containing substituents of phthalimidomethyl groups in number of 1 to 5, and the like.
 - [0085] Examples of yellow coloring pigments are C.I. Pigment Yellow 1, 2, 3, 4, 5, 6, 7, 10, 11, 12, 13, 14, 15, 16, 17, 23, 65, 73, 83, C.I. Vat Yellow 1, 3, 20 and the like.
- [0086] The above-described dyes and pigments may be used solely or may be used while being optionally mixed with one another to obtain desired hue of the toner. Incidentally, taking the environmental preservation and the sately to human being into consideration, a variety of edible coloring elements may preferably be used. The content of the coloring agents in the toner may widely altered depending on the desired coloration effects. Generally, in order to obtain the best toner properties, that is, in consideration of the printing coloration capability, the toner shape stability, and toner lesp, these coloring agents are used at a ratio in the range of from 0.1 to 80 parts by weight, preferably 0.5 to 20 parts by weight with the spect to 100 parts by weight of the binder resin.

Other components of toner

[0887] In the electrostatic latent image developing toner of the present invention may contain the following compounds other than the foregoing binder resin and the coloring agent components, to an extent (within a ratio less than the content of the binder resin) in which no undesired effect is caused in the present invention. Examples of such compounds include silicone resin; polypester; polyurethane; polyamide; epoxy resin; poly(inyl) butyral); resin; modified resin; terpene resin; phenolic resin; alightation or alleyclic hydrocarbor resin such as lower molecular weight polypropylene; aromatic type petroleum resin; and chlorinated paraffin and paraffin waxes. Among them, preferable waxes to be used an expractically lower molecular weight polypropylene; and ster type wax and aliphatic derivatives. Among them, preferable waxes to be used an expractically lower molecular weight polypropylene and its byproducts, lower molecular weight oply polyester, and ester type wax and aliphatic derivatives. Among these waxes, waxes separated based on the molecular weight of the waxes by various methods are also preferably used in the present invention. Further, after separation, the waxes may be modified to control the acid values, block-copolymerized, or graft-modified. [088] Specially, in the electrostatic latent image developing toner of the present invention, in the case such wax components as described above are added and these wax components are found practically dispersed in the binder resin in spherical and/or elliptical island state by cross-sectional observation of the toner by a transmission electron microscope, the toner is provided with excellent properties.

Method of producing toners

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(2088) Any conventionally known method may be employed for a practical method for producing an electrostatic latent image developing toner of the present invention having the constitution as described above. The electrostatic latent image developing toner of the present invention can be produced, for example, by a so-called pulverization method for obtaining a toner through the following steps. Specifically, the compound presented in the present invention described previously, reali materials such as binder rosis, and a wax to be added as necessary are sufficiently with one such as the she shall register, and a such as healing rolls, a kneader, an extruder and the like to make the resin material compatible with one another, and as cotoring agents, pigments, dyes, or magnetic materials and also additives such as metal compounds to be added as necessary are dispersed or dissolved in the resulting mixture, and after sollification of the mixture of present of the control of the control of the present invention with a decided particle size. In the above-described classification step, from an aspect of productivity, a multi-step classification apparatus is an enterably used.

[0090] In addition, the electrostatic latent image developing toner of the present invention with a desired particle size can be obtained by mixing and stifring the binder resin and the compound of the present invention in a solvent (e.g., aromatic hydrocarbons such as clotuene, xylene and the like, halogen compounds such as clotroform, eitylene dichloride, and the like, ketones such as acetoroe, methyl othyl ketone, and the like; amides such as dimethylformamide and the like), and then adding the resulting mixture to water to re-precipitate the solid, then filtering and drying the solid, and further pulvetizing it by a pulverizing apparatus such as a jet mill, a ball mill and the like, and finally classifying the pulvetized matter. In the above-described classification step, from an aspect of productivity, a multi-step classification apparatus is preferably used.

[0091] In addition, the electrostatic latent image developing toner of the present invention can be produced by a socalled polymerization method as follows. That is, in this case, the compound of the present invention, a polymerization monmer, and as coloring agents, pigments, dyes, or magnetic materials and also based on the necessity, additives such as a cross-infising agent, a polymerization initiator, waxes, and others are mixed and dispersed and in the presence of a surfactant or the like, the mixture is subjected to suspension polymerization to obtain a polymerization and coloring resin particle, and after the obtained particle is separated by solid-liquid separation, the particle is dried and classified incessary to obtain an electrostatic latent image developing toner of the present invention with a desired particle size. [0992] Furthermore, a coloring fine particle containing no charge controlling agent is produced by the above-described manner and then either solely or together with an externally added agent such as colloidad slice, the company presented in the present invention may be stuck and added to the surface of the particle by a mechanochemical method or the like.

(Externally added silica agent)

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[0093] In the present invention, a silica fine powder is preferably added externally to the toner produced in a manner as described above for improving the charge stability, development characteristic, fluidity and durability. The silica fine powder to be employed in this case can provide desirable effects if it has a specific surface area equal to or larger than 20 m²/g or higher (especially 50 to 400 m²/g) measured based on the nitrogen adsorption by BET method. The content

of the silica fine powder to be added is preferably 0.01 to 8 parts by weight, more preferably 0.1 to 5 parts by weight, with respect to 100 parts by weight of the toner particle. In this case, based on the necessity, the silica fine powder to be used in the case is preferably treated for the purpose of controlling the hydrophobility and charging property with silicone varnish, variously modified silicone varnish, variously modified silicone varnish, silicone oil, variously modified silicone oil, a siliane coupling agent, a siliane coupling agent having a functional group, and other organosilicon compounds. These treatment agent may be used by miking.

Inorganic powder

0094] Further, in order to improve the development capability and the durability, the following inorganic powder is preferably added. Examples of the powder are oxides of metals such as magnesium, zinc, aluminum, cerium, cobalt, iron, zirconium, chromium, manganese, strontium, tin, antimony and the like; compounded metal oxides such as calcium titanate, metal salts such as calcium carbonate, magnesium carbonate and aluminum carbonates; clay minerals such as kalein; posphate compounds such as apatie; silicon compounds such as salties; silicon compounds such as patie; silicon compounds such as carbonate; clay minerals such as kalein; posphate compounds such as patie; silicon compounds such as patie; silicon compounds so described, and silicon nitride; and carbon powder such as carbon black and graphite. Among them, fine powders of zinc oxide, aluminum oxide, cobalt oxide, manganese dioxide, strontium titanate, and magnesium titanate are preferably used.

Lubricant

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[0095] Further, the following lubricant powder may be added to the toner. For example, fluoro resin such as Teflon, poly(vinylidene fluoride) and the like; fluoride compounds such as carbon fluoride; aliphatic acid metal salts such as zince steartar; alliphatic acid derivatives such as aliphatic acid, aliphatic acid osters and the like; and molybdenum sulfide.

25 (Carrier)

[0096] The electrostatic latent image developing toner of the present invention having the above-described constitution is usable for a variety of conventionally known toners, solely as a non-magnetic mono-component developer, as a magnetic toner to be used as logically an anon-magnetic toner to be used as the magnetic carrier for composing a magnetic two-component developer, as a magnetic toner to be used solely for a magnetic mono-component toner. In this case, as the carrier to be used in the case of the two-component day promount ender or arrier may be used. More particularly, particles of surface-oxidized or non-oxidized metals such as iron, nickel, cobalt, manganese, chromium and rare earth metals, their alloys and oxides and having an average particle size of 20 to 300 µm may be used as the carrier particle. Further, the carrier to be used in the present invention are preferably the above-described carrier particle. Further, the carrier to be used in the present invention are preferably the above-described carrier particle whose surface bears or is coated with a substance such as styrene-based resin, acrylic resin, silicone resin, fluoro resin, polyester resin and the like

(Magnetic toner)

- 40 [0987] The electrostatic latent image developing toner of the present invention may be a magnetic toner by adding a magnetic material to be incore particle. In this case, the magnetic material may take a role also as a coloring agent. The magnetic material not be used in this case may be iron oxides such as magnetile, hematite and ferrite; metals such as iron, cobait and nickel; alloys of these metals with metals such as aluminum, oball, copper, lead, magnesium, in, zinc, aritimony, benyllium, bismuth, caditum, calcium, magneses, selenium, titanium, tungsten and variatium, and their mixtures. The magnetic material to be used in the present invention has an avertage particle size preferably 20 to 200 parts by weight to 100 parts by weight of the binder resin.
 - [0088] In addition, in order to give high image quality, it is required to precisely develop very small latent image dots and for this purpose, for example, it is preferable that the weight average particle size of the electrostatic latent mage developing toner of the present invention is controlled so that it is in the range of from 4 to 9 µm. That is, if the toner particle has a weight average particle size smaller than 4 µm, the transfer efficiency is decreased and a large amount of the transfer residual toner tends to remain on a photoconductor to result in an undesirable cause of uneven and irregular image formation attributed to fogging and transfer failures. Whereas, if the toner particle has a weight average 5 particle size larger than 9 µm, letters and the images tond to be oliminated.

[0099] In the present invention, the average particle size and the particle size distribution of the toner are measured by using Coulter Counter TA-II model or Coulter Multisizer (manufactured by Coulter Co.) or the like to which an interface (manufactured by Nikka Machine Co.) for outputting the distribution by number, the distribution by volume and a PC9801

personal computer (manufactured by NEC) are connected. As an electrolytic solution to be used at that time, an aqueous 1% NACI solution is prepared using first-grade sodium chloride. As the electrolytic solution, for example, a commercialized ISOTON R-II (produced by Coulter Scientific Japan Co.) may also be usable. A practical measurement method involves steps of adding 0.1 to 5 ml of a surfactant (preferably an alkybenzenesulfonic acid salt is used) as a dispersant to 100 to 150 ml of the above-described aqueous solution, further adding 2 to 20 mg of a sample to the resulting solution to obtain a specimen to be measured. At the time of measurement, the electrolytic solution in which the specimen to be measured is suspended is treated for dispersion for 1 to 3 minutes by an ultrascoil dispersing apparatus and then the volume and the number of the toner particles of 2 µm or larger are measured by the foregoing Coulter Counter TA-II model using 100 µm apertures as apertures and the distribution by volume and the distribution by number are calculated. Then, the weight average particle size (D4) on the bases of the volume calculated from the distribution by volume according to the present invention and the length average particle size (D1) on the bases of the number calculated from the distribution by number are acclusted from the distribution by wolume and the distribution by wolume and the distribution by wolume and the distribution by wolume according to the present invention and the length average particle size (D1) on the bases of the

(Charge level)

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[0100] In addition, the charge level of the electrostatic latent image developing toner of the present invention is preferably in the range of from -10 to -80 μC/g, more preferably from -15 to -70 μC/g per unit weight (two-component method) in improving the transfer efficiency in a transfer method using a transfer member with a voltage applied thereto. [0101] The method of measuring an charge level (a two-component tribo) by the two-component method employed in the present invention will be described as follows. A charge level measuring apparatus illustrated in FIG. 8 is used for the measurement. At first, under a specified environment, EFV 200/300 (produced by Powder Tec Co.) is used as a carrier and a bottle made of a polyethylene with a capacity of 50 to 100 ml is charged with a mixture of 9.5 g of the carrier and 0.5 g of a toner, an object to be measured, set in a shaking apparatus so controlled as to keep the amplitude constant, and shaken for a prescribed period in the shaking conditions of an amplitude of 100 mm and a shaking speed of 100 time reciprocation per 1 minute. Then, 1.0 to 1.2 g of the above mixture is placed in a measurement container 42 made of metal having a 500-mesh screen 43, and the measurement container 42 is covered with a metal lid 44 in the bottom of the charge level measuring apparatus shown in FIG. 8. The total weight of the measurement container 42 at that time is measured and determined as W1 (g). Next, the gas in the container is aspirated through a suction port 47 by an unillustrated aspirator (at least the portion contacting the measurement container 42 is made of an insulator) and an air ventilation adjustment valve 46 is controlled to control the pressure of the vacuum meter 45 to be 2.450 Pa (250 mmAq). Under such a state, aspiration is carried out for 1 minute to suck and remove the toner. The potential of a potentiometer 49 at that time is denoted as V (volt). The reference numeral 48 denotes a capacitor and the capacity is denoted as C (μ F). The weight of the entire measurement container after the aspiration is weighed and denoted as W2 (g). The friction charge level (µC/g) of the toner can be calculated according to the following equation from these measurement values.

Friction charge level(μ C/g) = C × V / (W1 - W2)

40 (Molecular weight distribution of binder resin)

[0102] The binder resin for use in the constituent material of the electrostatic latent image developing toner of the present invention preferably has a peak within the range of from 3,000 to 15,000 in a tow molecular weight region of the molecular weight distribution measured by GPC, especially, in the case of production by the pulverization method. That is, if the GPC peak exceeds 15,000 in the low molecular weight region, it sometimes becomes difficult to obtain a toner with a sufficiently improved transfer efficiency. Whereas if binder resin having a GPC peak of less than 3,000 is used, melting takes place easily at the time of surface treatment and therefore it is undestrable.

[0103] In the present invention, the molecular weight of the binder resin is measured by GPC (ge) permeation chromatography. A practical GPC measurement method is carried out as follows: a toner previously extracted with THF (letrahydrofuran) solvent for 20 hours using a Soxhlet extractor is used as a sample for measurement and using columns to A801, 802, 803, 804, 805, 806, and 807 manufactured by Showa Derko KK. and calibration curves of standardized polystyrene resins, the molecular weight distribution is measured. Further, in the present invention, it is preferable that the binder resin with the ratio (Mw/Mn) of the weight average molecular weight (Mw) to the number-average molecular weight (Mm) measured as described above being in the range of form 2 to 100 is used.

55 (Glass transition temperature of toner)

[0104] Further, the toner of the present invention is preferably adjusted by using a proper material so as to have a glass transition temperature Tg in the range of from 40 to 75°C, more preferably 52 to 70°C, from a viewpoint of fixation

and storage stability. In this case, the measurement of the glass transition temperature Tg may be carried out using a high precision and internally heating input compensation type differential scanning calorimeter, for example, DSC-7 manufactured by Perkin Elmer Co, may be employed. The measurement method is carried out according to ASTM D3416-82. In the present invention, in the case of measuring the glass transition temperature Tg, it is preferable that a measurement sample is once heated to cancel the entire hysteresis and then quenched and again heated at a heating rate of 10°C/min to employ the DSC curve measured during the healing form 0 to 200°C.

(Image formation method)

- 0 [015] The electrostatic latent image developing toner of the present invention having the configuration described above is particularly preferably applied to an image formation method comprising at least an charging step of applying a voltage to a charge member from the outside to charge an electrostatic latent image carrier, a step of forming an electrostatic latent image on the charged electrostatic latent image carrier, a development step of developing the electrostatic latent image or the toner to form a toner image on the electrostatic latent image carrier, a transfer siner, a transfer siner of transferring the toner image on the electrostatic latent image carrier to an object recording material, and a heat-fixation step of heat-fixing the toner image on the object recording material, or a image formation method with the transfer step consisting of a first transfer step of transferring the toner image on the intermediate transfer body to the object recording material.
- 20 [1016] The present invention provides an innovative polyhydroxyalkaneate with a sulfonic group as a hydrophilic group and its derivative introduced therein and a method of producing the same. In this way, the innovative polyhydroxyalkaneate is accellent in melt-processability, and also excellent in biocompatibility owing to its hydrophilic nature, and can thus be excelled to be apolled as medical flexible members and the like.
 - [0107] In addition, as described above and below, according to the present invention, addition of one or more types of compounds presented in the present invention to an electrostatic latent image developing toner composition as a charge controlling agent makes it possible to provide an electrostatic latent image developing toner having an excellent electrifiability, improving the dispersibility of the compound in the toner resin and the spent characteristic thereof, causing no image fog even when the image is outputed in the image forming apparatis, and being sexcellent in transfersibility and highly applicable to an electrophotographic process. In addition, because the charge controlling agent for use in the present invention is coloriess or only weakly colored, any coloring agent can be selected according to the color required for the color toner, and the original color of a dye or pigment is not hindered.

(Examples)

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- [0108] The present invention will be described further in detail below with reference to Examples, although the method of the present invention should not be limited to the Examples.
 - [0109] First, the method of producing sodium 2-(2'-mercaptoethyl)amide-2-methylpropanesulfonate of formula (24)

as one of compounds each expressed by chemical formula (19) will be described.

[0110] 15.2 g of 2-acrylamide-2-methylpropanesulfonic acid, 2.8 g of sodium hydrate and 0.01 g of benzoyl peroxide were dissolved in 48.5 g of medhanol at a room temperature, and gas in the system was replaced with nitrogen. Then, 26.7 g of thioacetic acid was added while maintaining the temperature in the system at 15 to 19°C, and the mixture was thereafter left under reflux for 4 hours, holding the system at the temperature of 45 to 80°C. After cooling, 600 g of isopropy lefter was added to wash the mixture. The insoluble matter was dired so that the velight thereof was reduced to 22 g. The obtained insoluble matter was dissolved in 66 g of methanol at a room temperature, and gas in the system was replaced with nitrogen, 0.7 g of so double mydrate was added, and was stirred for 3 hours with the temperature in the system being kept at 39 to 41°C. After cooling, 1.2 g of accide acid was added, and fiveration the solvent was distilled away to obtain thiolated 2-acrylamide-2-methylpropanesulfonic acid. Furthermore, the 1H-NMR sectrum was used to ensure that the thiolation had been done in a quantitative manner.

[0111] The following Examples 1 to 4 are examples of methods of producing the polyhydroxyalkanoate of the present invention using as a raw material 2-(2'-mercaptoethyl)amide-2-methylpropanesulfonate obtained in the above manner and the polyhydroxyalkanoate produced by the method of production by microorganism described previously. However, the polyhydroxyalkanoate and the method of producing the same should not be solely dependent on the above raw materials.

Example 1

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[0112] Method (1) of producing polyhydroxyalkanoate containing units expressed by chemical formulas (25) and (26)

[0113] 900 mg of polyhydroxyalkanoate (average molecular weight: Mn=36000, Mw=66000 (measured by gel permeation chromatography (GPC); Tosoh HLC-8220, column; Tosoh TSK-GEL SuperHM-H, solvent; chloroform, polystyrene equivalent)) containing 9.5 mol% in total of 3-hydroxy-8-bromooctanoic acid unit and 3-hydroxy-6-bromohexanoic acid unit, 89.1 mol% of 3-hydroxy-5-phenylvaleric acid and 1.4 mol% of other components (straight-chain 3-hydroxyalkanoic acid having 4 to 12 carbon atoms and straight-chain 3-hydroxyalka-5-enoic acid having 10 or 12 carbon atoms) was dissolved in 12 ml of N,N-dimethylformamide at a room temperature, and gas in the system was replaced with nitrogen. Then, while holding the system at a room temperature, 825 mg of sodium 2-(2'-mercaptoethyl)amide-2-methylpropanesulfonate dissolved in 18 ml of N.N-dimethylformamide was added, and 330 ul of diethylamine was further added, and was stirred at a room temperature for 24 hours.

[0114] After the reaction was completed, the mixture was put in 300 ml of diethyl ether to allow reprecipitation for removing N,N-dimethylformamide as a reaction liquid. The resulting precipitate was collected by centrifugal separation. [0115] This precipitate was put in 300 ml of water, stirred and washed. The precipitate obtained at this time was collected by carrying out centrifugal separation. This precipitate was dried under reduced pressure to obtain 244 mg of polyhydroxyalkanoate.

[0116] This PHA was analyzed using nuclear magnetic resonance apparatus under the following conditions. Measuring apparatus

FT-NMR: Bruker DPX400

resonance frequency: 1H=400MHz

Measuring apparatus nuclear species to be measured: 1H

solvent used: DMSO-d6

reference: capillary-encapsulated DMSO-d6

measuring temperature: 40°C

The 1H-NMR spectrum chart is shown in FIG. 1, and the composition of PHA calculated from the results obtained by the ¹H-NMR spectrum is shown in Table 1.

Table 1

	Content of each unit (mol%)
Chemical formulas (25) and (26)	4.8
Chemical formula (27)	93.0
Other polyhydroxyalkanoate	2.2

From the results, this polyhydroxyalkanoate was found to contain 4.8 mol% in total of units expressed by chemical formulas (25) and (26) and 93.0 mol% of 3-hydroxy-5-phenylvaleric acid unit expressed by chemical formula

[0118] The molecular weight of the obtained PHA was measured by gel permeation chromatography (GPC; Tosoh HLC-8220, column: Tosoh TSK-GEL SuperHM-H, solvent; chloroform, polystyrene equivalent) and found as follows: Mn=10000 and Mn=194000.

Example 2

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[0119] Method (2) of producing polyhydroxyalkanoate containing units expressed by chemical formulas (25) and (26) [0120] 869 mg of polyhydroxyalkanoate (severage molecular weight: Ma-38040, Mwei65200 measured by the gel permeation chromatography described in Example 1) containing 7.4 mol% in total of 3-hydroxy-8-bromoclasnoic acid unit and 3-hydroxy-8-bromoclasnoic acid unit and 3-hydroxy-8-bromoclasnoic acid (8.71 mol% of 3-hydroxy-9-bromoclasnoic acid and 5.5 mol% of other components (straight-chain 3-hydroxyalkanoic acid having 4 to 12 carbon atoms and straight-chain 3-hydroxyalka-6-andioc acid having 10 or 12 carbon atoms was dissolved in 9 ml of N,N-dimethylformamide at a room temperature, and gas in the system was replaced with introgen. Then, while holding the system at a room temperature, along 2-(2-mercaptoethyl)amide-2-methylpropanesulfonate dissolved in 16 ml of N,N-dimethylformamide was added, and 208 µJ of dishtylamine was then added, and was stirred at a room temperature (2-4 hours.

[0121] After the reaction was completed, N.N-dimethyformamide as a reaction liquid was once distilled away by a rotary veoporator, and the mixture was again dissolved in 3 ml of N.N-dimethyformamide, and was thereafter put in 300 ml of pure water to allow reprecipitation. The resulting precipitate was collected by centrifugal separation. This precipitate was suspended again with 50 ml of pure water, and subjected to centrifugal separation to collect the precipitate for washing. This washing operation was conducted three times, followed by drying the precipitate under reduced pressure to obtain 619 mg of polyhydroxyalkanoate.

[0122] For this PHA, measurements were carried out using the nuclear magnetic resonance apparatus under the same condition as Example 1.

[0123] The composition of PHA calculated from the results obtained by the ¹H-NMR spectrum is shown in Table 2.

Table 2

	Content of each unit (mol%)
Chemical formulas (25) and (26)	4.0
Chemical formula (28)	91.0
Other polyhydroxyalkanoate	5.0

[0124] From the results, this polyhydroxyalkanoate was found to contain 4.0 mol% in total of units expressed by chemical formulas (25) and (26) and 91.0 mol% of 3-hydroxy-5-phenoxyvaleric acid unit expressed by chemical formula (28)

[0125] The molecular weight of the obtained PHA was measured by the method described in Example 1 using gel permeation chromatography and found as follows: Mn=3100 and Mw=9100.

Example 3

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[0126] Method (1) of producing polyhydroxyalkanoate containing units expressed by chemical formulas (29), (30) and (31)

[0127] 449 mg of polyhydroxyalkanoate (average molecular weight: Mn=48000, Mw=111000 measured by the gel permeation chromatography described in Example 1) containing 7.1 mol% in total of 3-hydroxy-1-brownoundocanoid acid unit, 3-hydroxy-9-brownonanoia caid unit and 3-hydroxy-7-brownoheptanoic acid, 79.3 mol% of 3-hydroxy-5-phe-

noxyvaleric acid and 13.7 mol% of other components (straight-chain 3-hydroxyalkanoic acid having 4 to 12 carbon atoms and straight-chain 3-hydroxyalka-5-enoic acid having 10 or 12 carbon atoms) was dissolved in 6 mi of N,N-dimethylformanide at a room temperature, and gas in the system was replaced with introgen. Then, while holding the system at a room temperature, 265 mg of sodium 2-(2-mercaptoethyl)amide-2-methylpropanesulfonate dissolved in 8 ml of N,N-dimethylformamide was added, and 106 µl of diethylamine was then added, and was stirred at a room temperature for 24 hours.

[0128] After the reaction was completed, NN-dimbtyformamide as n-direction liquid was once distilled away by a ordary evaporator, and the mixture was again dissolved in 2 ml of NN-dimbtyformamide, and was thereafter put in 200 ml of pure water to allow reprecipitation. The resulting precipitate was collected by centrifugal separation. This precipitate was suspended again with 50 ml of pure water, and subjected to centrifugal separation to collect the pre-cipitate is usepended again with 50 ml of pure water, and subjected to centrifugal separation to collect the pre-cipitate for washing. This washing operation was conducted three times, followed by drying the precipitate under reduced pressure to obtain 389 mo of polyhydrovaytikanoate.

[0129] For this PHA, measurements were carried out using the nuclear magnetic resonance apparatus under the same condition as Example 1.

[0130] The composition of PHA calculated from the results obtained by the ¹H-NMR spectrum is shown in Table 3.

Table 3	
	Content of each unit (mol%)
Chemical formulas (29), (30) and (31)	2.1
Chemical formula (28)	85.3
Other polyhydroxyalkanoate	12.6

9 14, C CH, SO, Na PH, C CH, SO, Na PH,

[0131] From the results, this polyhydroxyalkanoate was found to contain 2.1 mol% in total of units expressed by chemical formulas (29), (30) and (31) and 85.3 mol% of 3-hydroxy-5-phenoxyvaleric acid unit expressed by chemical formula (28).

[0132] The molecular weight of the obtained PHA was measured by the method described in Example 1 using gel permeation chromatography and found as follows: Mn=3400 and Mw=14500.

Example 4

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[0133] Method (2) of producing polyhydroxyalkanoate containing units expressed by chemical formulas (29), (30) and (31)

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[0134] 300 mg of polyhydroxyalkanoate containing 22.0 mol% in total of 3-hydroxy-11-bromoundacanoic acid unit, 3-hydroxy-9-bromonanoic acid unit, and 3-hydroxy-7-bromohapanoic acid. 80, 0 mol% of 3-hydroxy-5-(phenysiultony) valeric acid and 9.0 mol% of 5-hydroxy-5-bromohapanoic acid. 80 mol% of 12 carbon atoms and straight-of-ain 3-hydroxyalkanoic acid having 4 to 12 carbon atoms and straight-of-ain 3-hydroxyalkanoic acid having 4 to 12 carbon atoms was dissolved in 14 mil of N-4-dimethyl-formamide at a room temperature, and gas in the system was replaced with nitrogen. Then, while holding the system at a room temperature, 446 mg of sodium 2-(2'-mercaptoethyl)amide-2-methylpropenesulfonate dissolved in 6 mil of N-1 dimethylformamide was added, and 180 µ of dishylamine was then added, and was stirred at a room temperature.

[0135] After the reaction was completed, N,N-dimethylformamide as a reaction liquid was once distilled away by a rotary evaporator, and the mixture was again dissolved in 1 2 ml of N,N-dimethylformamide, and was thereafter put in 200 ml of pure water to allow reprecipitation. The resulting precipitate was collected by centrifugal separation. This precipitate was suspended again with 50 ml of pure water, and subjected to centrifugal separation to collect the precipitate for washing. This washing operation was conducted three times, followed by drying the precipitate under reduced pressure to obtain 227 mg of polyhydroxyalkanoate.

[0136] For this PHA, measurements were carried out using the nuclear magnetic resonance apparatus under the same condition as Example 1.

[0137] The composition of PHA calculated from the results obtained by the ¹H-NMR spectrum is shown in Table 4.

Table 4

	Content of each unit (mol%)
Chemical formulas (29), (30) and (31)	2.7
Chemical formula (32)	70.6
Other polyhydroxyalkanoate	26.7

[0138] From the results, this polyhydroxyalkanoate was found to contain 2.7 mol% in total of units expressed by chemical formulas (29), (30) and (31) and 70.6 mol% of 3-hydroxy-5-(phenylsulfonyl)valeric acid unit expressed by chemical formula (32).

[0139] The PHA obtained in Examples 1 to 3 described above, which was classified into exemplary compounds (1) to (3) as shown in Table 5, was used as a charge controlling agent to produce various kinds of toners, and evaluations were carried out (Examples 5 to 44).

Table 5

Exemplary compound (1)	PHA of Example 1
Exemplary compound (2)	PHA of Example 2
Exemplary compound (3)	PHA of Example 3

Example 5

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[0140] First, an aqueous Na₆PO₄ solution was added in a 2 liter four-necks flask equipped with a high-speed stirring apparatus TK-Homomiker, and was heated at 60°C with the number of rotations kept at 10,000 rpm. An aqueous CaCl₂ solution was slowly added therein to prepare an aqueous dispersing medium containing a very small water slightly solutile dispersant Ca₆(PO₂).

[0141] On the other hand, the following compositions were dispersed for 3 hours using a ball mill, followed by adding therein 10 parts by weight of release agent (ester wax) and 10 parts by weight of 2,2*-azobis(2,4-dimethylvaleronitrile) as a polymerization initiator to prepare a polymerizable monomer composition.

- styrene monomer 82 parts by weight
- ethylhexyl acrylate monomer 18 parts by weight
- divinylbenzene monomer 0.1 parts by weight
- cyan coloring agent (C.I. Pigment Blue 15)
 6 parts by weight
- oxidized polyethylene resin (molecular weight 3200, acid number 8)
 5 parts by weight
 - exemplary compound (1) 2 parts by weight

[0142] Then, the polymerizable monomer composition obtained as described above was put in the aqueous dispersing medium prepared previously to carry out the granulation with the number of rotations being kept at 10,000 rpm. Thereafter, the composition was made to undergo a reaction at 65°C for 3 hours white being stirred with a patidle stirring blade, and was thereafter polymerized at 80°C for 6 hours to complete the polymerization reaction. After the reaction was completed, the suspension was cooled, and an acid was added therein to dissolve the water slightly soluble dispersant $Ca_0(PO)_{3/2}$, followed by filtering, nirsing and drying the solution to obtain blue polymerized particles (1). The particle size of the obtained blue polymerized particles (1) measured using Coulter Counter Multilazer (manufactured by Coulter Co.) was 7.1 μ m as a weight average particle size, and the ratio of lines (the abundance ratio of particles with the size 40.3.1 μ m or smaller in the number distribution) was 5.5% by number.

[0143] As a fluidity improver, 1.3 parts by weight of hydrophobic silica fine powder (BET: 270 m²/g) treated with hexamethyl disilazane were externally added to 100 parts by weight of blue polymerized particles(1) prepared as

described above through dry-mixing by a Henshel mixer, whereby a blue toner (1) of this Example was obtained. In addition, 7 parts by weight of blue toner (1) were mixed with 83 parts by weight of resin-coated magnetic ferrite carrier (average particle size: 45 µm) to prepare a two-component type blue developer (1) for magnetic bursh development.

Examples 6 and 7

[0144] Blue toners (2) and (3) of Examples 6 and 7 were obtained in the same manner as Example 5 except that 2.0 part by weight of exemplary compounds (2) and (3) are used, respectively, in place of exemplary compound (1). In addition, two-component type blue developers (2) and (3) were obtained, respectively. In the same manner as Example 5 using the blue toners (2) and (3) and the above described resin-coated magnetic ferrite carrier. For these blue toners (2) and (3) as well as the two-component type blue developers (2) and (3), the properties of toners were measured in the same manner as Example 5. and the results thereof are shown in Table 6.

Comparative Example 1

[0145] A blue toner (4) of Comparative Example 1 was obtained in the same manner as Example 5 except that no charge controlling agent was used. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Table 6. In addition, a two-component type blue developer (4) of Comparative Example 1 was obtained in the same manner as Example 5 using this toner.

Evaluation

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[0146] For the two-component type blue developers (1) to (3) obtained in the Examples 5 to 7 and the two-component type blue developer (4) obtained in the Comparative Example 1, the charge levels of toners after stirring for 10 and 300 seconds were measured under conditions of normal temperature and normal humidity (26°C, 60% RH) and high temperature and high humidity (30°C, 80% RH) and high temperature and high humidity (30°C, 80% RH) and high temperature and high humidity (30°C, 80% RH) and high temperature and high humidity (30°C, 80% RH) using the previously described method of measuring charge levels. Then, measurement values of two-component blow-off charge levels were rounded off to the first decimal place, and the resultant values were evaluated according to the following ordinar. The results are shown together in Table 6.

30 Electrifiability

[0147]

- A: Excellent (-20 μC/g or lower)
- B: Good (-19.9 to -10.0 uC/a)
- C: Usable (-9.9 to -5.0 µC/a)
- D: Unusable (-4.9 µC/g or higher)

Table C

40	Table 6									
	Examples	Toners Number: Blue			Electrifiability					
45			Weight average particle size (µm)	Ratio of fines (% by number)	Normal temperature and normal humidity (Q/M)		High temperature and high humidity (Q/M)			
50					Stirring for 10 seconds	Stirring for 300 seconds	Stirring for 10 seconds	Stirring for 300 seconds		
	5	1	7.1	5.5	А	А	A	Α		
	6	2	7.2	5.3	А	А	А	Α		
55	7	3	7.0	5.1	В	А	В	Α		

Table 6 (continued)

5	Examples	Toners Number: Blue	Particle size	distribution	Electrifiability	•				
			Weight average particle size (µm)	Ratio of fines (% by number)	Normal temperature and normal humidity (Q/M)		High temperature and high humidity (Q/M)			
					Stirring for 10 seconds Stirring for 300 seconds		Stirring for 10 seconds	Stirring for 300 seconds		
15	Comparative Example 1	4	7.0	5.2	D	D	D	D		

Examples 8 to 10

- [0148] Yellow toners (1) to (3) were obtained in the same manner as Example 5 except that 2 0 parts by weight of exemplary compounds (1) to (3) were used, and a yellow coloring agent (Hansa yellow G) was used in place of the cyan coloring agent. The properties of these toners were measured in the same manner as Example 5, and the results thereof are shown in Table 7. In addition, two-component type yellow developers (1) to (3) were obtained in the same manner as
- 25 Example 5 using these, toners.

Comparative Example 2

- [0149] A yellow toner (4) of Comparative Example 2 was obtained in the same manner as Example 5 except that no charge controlling agent was used, and that the yellow coloring agent (Hansa yellow G) was used in place of the cyan coloring agent. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Tablo 7. In addition, a two-component type yellow developer (4) of Comparative Example 2 was obtained in the same manner as Example 6 using this toner. Evaluation
- [0150] For the two-component type yellow developers (1) to (3) obtained in the Examples 8 to 10 and the two-component type yellow developer (4) obtained in the Comparative Example 2, the charge levels of toners after stirring for 10 and 300 seconds were measured under conditions of normal temperature and normal humidity (26°C, 60% RH) and high temperature and high humidity (30°C, 80% RH) using the previously described method of measuring charge levels.
- [0151] Then, measurement values of two-component blow-off charge levels were rounded off to the first decimal place, and the resultant values were evaluated according to the following criteria. The results are shown together in Table 7. Electrifiability
 - A: Excellent (-20 µC/g or lower)
 - B: Good (-19.9 to -10.0 μC/g)

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- C: Usable (-9.9 to -5.0 µC/g)
- D: Unusable (-4.9 µC/g or higher)

Table 7

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5	Examples	Toners Number: Yellow	Particle size	rticle size distribution Electriflability				
10			Weight average particle size (µm)	Ratio of fines (% by number)			High tempera humidity (Q/N	
					Stirring for 10 seconds	Stirring for 300 seconds	Stirring for 10 seconds	Stirring for 300 seconds
15	8	1	7.0	5.4	Α	Α	Α	Α
	9	2	7.2	5.3	Α	Α	Α	Α
	10	3	6.9	5.5	В	Α	В	В
20	Comparative Example 2	4	7.2	4.9	D	D	D	D

Examples 11 to 13

[0152] Black toners (1) to (3) of were obtained in the same manner as Example 5 using 2.0 parts by weight of exemplary compounds (1) to (3) except that a carbon black (OBP oil absorption 110 mU/100 g) was used in place of the cyan coloring agent. The properties of these toners were measured in the same manner as Example 5, and the results thereof are shown in Table 8. In addition, two-component type black developers (1) to (3) were obtained in the same manner as Example 5 using these toners.

30 Comparative Example 3

[0153] A black toner (4) of Comparative Example 3 was obtained in the same manner as Example 5 except that no charge controlling agent was used and that the carbon black (DBP oil absorption 110 m1/100 g) was used in place of the cyan coloring agent. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Table 8. In addition, a two-component type black developer (4) of Comparative Example 3 was obtained in the same manner as Example 6 using this flower.

Evaluation

[0154] For the two-component type black developers (1) to (3) obtained in the Examples 11 to 13 and the two-component type black developer (4) obtained in the Comparative Example 3, the charge levels of toners after stirring for 10 and 800 seconds were measured under conditions of normal temperature and normal hundidity (25°C, 60% RH) and high temperature and high hundidity (30°C, 80% RH) using the previously described method of measuring charge levels. Then, measurement values of two-component blow-off charge levels were rounded off to the first decimal place, and the results are shown together in Tables.

Electrifiability

[0155]

- A: Excellent (-20 μC/g or lower)
 - B: Good (-19.9 to -10.0 µC/a)
 - C: Usable (-9.9 to -5.0 µC/g)
- D: Unusable (-4.9 μC/g or higher)

Table 8

	11000								
5	Examples	Toners Number: Black	Particle size distribution Electrifiability						
10			Weight average fines (% by number) size (µm) Ratio of average rounder Normal temperature and normal humidity (Q/M) humidity (Q/M) humidity (Q/M) humidity (Q/M) humidity (Q/M) humidity (Q/M) humidity (Q/M)						
					Stirring for 10 seconds	Stirring for 300 seconds	Stirring for 10 seconds	Stirring for 300 seconds	
15	11	1	7.0	5.3	Α	Α	Α	А	
	12	2	7.1	5.4	Α	A	Α	А	
	13	3	6.9	5.2	В	В	В	В	
20	Comparative Example 3	4	6.9	5.3	D	С	D	С	

Example 14

[0156]

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- stylene-butylacrylate copolymer resin (glass transition temperature 70°C)
 100 parts by weight
- magenta pigment (C.I. Pigment Red 114)
 5 parts by weight
- exemplary compound (1)
 2 parts by weight

O157] The above described compositions were mixed and melt-kneaded by a blaxial extruder (UD=30). The resulting mixture was cooled and then roughly ground by a harmer mill, and finely being ground by a jet mill. The resultant powder was classified to obtain magenta coloring particles (1). The weight average particle size and the ratio of fines of the magenta coloring particles (1) were 7.0 µm and 5.2% by number.

[0158] As a fluidity improver, 1.5 parts by weight of hydrophobic silica fine powder (BET: 250 m²/g) treated with hexamethy disliazane were dry-mixed with 100 parts by weight of the magenta coloring particles (1) by a Henshel mixer, whereby a magenta toner (1) of this Example was obtained. In addition, 7 parts by weight of the resulting magenta toner (1) were mixed with 95 parts by weight of resin-coated magnetic fartie carrier (average particle size: 45 mit to proper a two-component type magenta developer (1) for magnetic brush development. The properties of this toner were measured in the same maner as Example 5, and the results thereof are shown in Table 3.

Examples 15 and 16

[0159] Magenta toners (2) and (3) of Examples 15 and 16 were obtained in the same manner as Example 14 except that 2.0 parts by weight of each of exemplary compounds (2) and (3) were used in place of exemplary compound (1). The properties of these toners were measured in the same manner as Example 5, and the results thereof are shown in Table 9. In addition, two-component type magenta developers (2) and (3) were obtained in the same manner as Example 14 using these toners.

Comparative Example 4

[0160] A magenta toner (4) of Comparative Example 4 was obtained in the same manner as Example 14 except that no charge controlling agent was used. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Table 9. In addition, a two-component type magenta developer (4) of Comparative Example 4 was obtained in the same manner as Example 14 using this toner.

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Evaluation

[0161] For the two-component type magenta developers (1) to (3) obtained in the Examples 14 to 16 and the two-component type magenta developer (4) obtained in the Comparative Example 4, the charge levels of toners after stirring for 10 and 300 seconds were measured under conditions of normal temperature and normal humidity (25°C, 60% RH) and high temperature and high humidity (30°C, 60% RH) using the previously described method of measuring charge levels. Then, measurement values of two-component blow-off charge levels were rounded off to the first decinal place, and the resultant values were evaluated according to the following criteria. The results are shown together in Table 9.

Electrifiability

[0162]

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A: Excellent (-20 μC/g or lower)

B: Good (-19.9 to -10.0 μC/g)

C: Usable (-9.9 to -5.0 µC/g)

D: Unusable (-4,9 µC/g or higher)

Table 9

	Table 9									
			Particle size	distribution	Electrifiability					
25	Examples	Toners Number: Red	Weight Ratio of average fines (% by number) size (µm)							
30					Stirring for 10 seconds	Stirring for 300 seconds	Stirring for 10 seconds	Stirring for 300 seconds		
	14	1	7.0	5.2	Α	Α	Α	Α		
	15	2	7.1	5.1	Α	Α	Α	Α		
35	16	3	6.9	5.3	В	Α	В	Α		
	Comparative Example 4	4	7.1	5.1	D	С	D	С		

Examples 17 to 19

[0163] Black toners (5) to (7) were obtained in the same manner as Example 14 using 2.0 parts by weight of examplary compounds (1) to (3) except that a carbon black (DBP oil absorption 110 mL/100 g) was used in place of the magenta pigment. The properties of these toners were measured in the same manner as Example 5, and the results thereof are shown in Table 10. In addition, two-component type black developers (5) to (7) were obtained in the same manner as Example 14 using these toners.

Comparative Example 5

[0164] A black toner (8) of Comparative Example 5 was obtained in the same manner as Example 14 except that no charge controlling agent was used and that the carbon black (OBP 6) ill absorption 110 mL/100 g) was used in place of the magenta pigment. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Table 10. In addition, a two-component type black developer (8) of Comparative Example 5 was obtained in the same manner as Example 14 surging this toner. Evaluation

[0165] For the two-component type black developers (6) to (7) obtained in the Examples 17 to 19 and the twocomponent type black developer (8) obtained in the Comparative Example 6, the charge levels of loners after stirring for 10 and 300 seconds were measured under conditions of normal temperature and normal humidity (26°C, 80% RH) and high temperature and high humidity (30°C, 80% RH) using the previously described method of measuring charge.

levels. Then, measurement values of two-component blow-off charge levels were rounded off to the first decimal place, and the resultant values were evaluated according to the following criteria. The results are shown together in Table 10.

Electrifiability

[0166]

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A: Excellent (-20 µC/g or lower)

B: Good (-19.9 to -10.0 µC/a)

C: Usable (-9.9 to -5.0 uC/g)

D: Unusable (-4.9 µC/g or higher)

				Tab	le 10			
15	Examples	Toners Number: Black	Particle size	distribution	Electrifiability			
20			Weight average particle size (µm) Ratio of fines (% by number)		Normal temp normal humid		High tempera humidity (Q/N	
25					Stirring for 10 seconds	Stirring for 300 seconds	Stirring for 10 seconds	Stirring for 300 seconds
	17	5	7.1	5.4	Α	Α	A	Α
	18	6	7.0	5.3	A	Α	A	Α
30	19	7	6.9	5.2	В	Α	В	В
30	Comparative Example 5	8	7.0	5.7	D	С	D	D

Example 20

[0167]

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- polyester resin 100 parts by weight
- carbon black (DBP absorption 110 ml/100 g) 5 parts by weight
- exemplary compound (1) 2 parts by weight

[0168] The polyester resin was synthesized as follows: 751 parts of bisphenol A propylene oxide 2 mol adduct, 104 parts of terephtalic acid and 167 parts of trimellitic anhydride were polycondensed with two parts of dibutyltin oxide as a catalyst to obtain a polyester resin having a softening point of 125°C.

[0169] The above described compositions were mixed and melt-kneaded by a biaxial extruder (L/D=30). The resulting mixture was cooled and then roughly ground by a hammer mill, and finely ground by a let mill. The resultant powder was classified to obtain black coloring particles (9). The weight average particle size and the ratio of fines of the black coloring particles (9) were 7.7 µm and 5.0% by number.

[0170] As a fluidity improver, 1.5 parts by weight of hydrophobic silica fine powder (BET: 250 m²/g) treated with hexamethyl disilazane were dry-mixed with 100 parts by weight of the black coloring particles (9) by a Henshel mixer, whereby a black toner (9) of this Example was obtained. In addition, 7 parts by weight of the resulting black toner (9) were mixed with 93 parts by weight of resin-coated magnetic ferrite carrier (average particle size; 45 µm) to prepare a two-component type black developer (9) for magnetic brush development. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Table 11.

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Examples 21 and 22

[0171] Black toners (10) and (11) of Examples 21 and 22 were obtained in the same manner as Example 20 except that 2.0 parts by weight of each of exemplary compounds (2) and (3) were used in place of exemplary compound (1). The properties of these toners were measured in the same manner as Example 5, and the results thereof are shown in Table 11. In addition, two-component type black developers (10) and (11) were obtained in the same manner as Example 20 using these toners.

Comparative Example 6

[0172] A black toner (12) of Comparative Example 6 was obtained in the same manner as Example 20 except that no exemplary compound (1) was used. The properties of this toner were measured in the same manner as Example 5, and the results thereof are shown in Table 11. In addition, a two-component type black developer (12) of Comparative Example 6 was obtained in the same manner as Example 20 using this toner. Evaluation

[0173] For the two-component type black developers (9) to (11) obtained in the Examples 20 to 22 and the twocomponent type black developer (12) obtained in the Comparative Example 6, the charge levels of toners after stirring for 10 and 300 seconds were measured under conditions of normal temperature and normal humidity (25°C, 60% RH) and high temperature and high humidity (30°C, 80% RH) using the previously described method of measuring charge levels. Then, measurement values of two-component blow-off charge levels were rounded off to the first decimal place, and the resultant values were evaluated according to the following criteria. The results are shown together in Table 11.

Electriflability

[0174]

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- A: Excellent (-20 µC/a or lower)
- B: Good (-19.9 to -10.0 μC/g)
- C: Usable (-9.9 to -5.0 µC/q)
- D: Unusable (-4.9 µC/g or higher)

				Tab	le 11				
35	Examples	Toners Number: Black	Particle size	distribution	Electrifiability				
40			Weight average particle size (µm)	Ratio of fines (% by number)	Normal temp normal humid		High tempera humidity (Q/N		
					Stirring for 10 seconds	Stirring for 300 seconds	Stirring for 10 seconds	Stirring for 300 seconds	
45	20	9	7.7	5.0	Α	Α	Α	Α	
	21	10	7.6	4.9	Α	Α	Α	Α	
	22	11	7.4	5.2	В.	В	В	Α	
50	Comparative Example 6	12	7.5	4.9	D	С	D	С	

Examples 23 to 40 and Comparative Examples 7 to 12

[0175] First, an image forming apparatus used in the image formation methods of Examples 23 to 40 and Comparative Examples 7 to 12 will be described. FIG. 2 is a schematic explanatory view of the cross section of an image forming apparatus for carrying out the image formation methods of Examples and Comparative Examples of the present invention. A photoconductor drum 1 shown in FIG. 2 has a photosensitive layer 1a having an organic photo semiconductor

FP 1 336 635 A1

on a substrate 1b, and is configured to rotate in the direction indicated by the arrow, and its surface is electrical charged at a potential of about -600 V by a charge roller 2 being a charge member situated opposite by high photoconductor drum 1 and contacting and rotating with the drum. As shown in FIG. 2, the charge roller 2 has a cored bar 2b covered with a conductive desilic layer 2a.

- 5 [0178] Next, the photoconductor drum 1 with its surface electrically charged is exposed to light 3 and at this time, oxidd operations are performed on the photoconductor by a polygon mirror according to digital larges information, whereby an electrostatic latent image with the potential of the exposed area being -100 V and the potential of the dark area being -500 V and the potential of the dark area being -500 V is formed. Subsequently, this electrostatic latent image on the photoconductor drum 1 is reversed eveloped and thereby actualized using a plurally of development apparatuses 4-1, 4-2, 4-3 and 4-4, and thus a toner of image is formed on the photoconductor drum 1. At this time, the two-component type developers obtained in Examples 5 to 22 and Comparative Examples 1 to 6 were individually used as developers to form a toner image with a yellow toner, a magenta toner, a cyan toner or a black toner. FIG. 3 is an enlarged sectional view of principal parts of development apparatuses 4 for two-component type developers used at that time.
- [0177] Then, the toner image on the photoconductor drum 1 is transferred to an intermediate transfer body 5 contacting and rotating with the photoconductor drum 1. As a result, a four-color cool combination developed image is formed on the intermediate transfer body 5. A non-transferred toner remaining on the photoconductor drum 1 without being transferred is collected in a container 9 for residual toners by a cleaner member 8.
- [0178]. The intermediate transfer body 5 is constituted by a cored bar 5b as a support and an elastic layer 5a provided thereon as shown in FIG. 2. In this Example, the intermediate body 5 having the cored bar 5b coated with the elastic layer 5b with a carbon black as a conductivity producer sufficiently dispersed in nitrile-butadene rubber (NBR) was used. The degree of hardness of the elastic layer 5b measured in accordance with "JIS K-6301" was 30 degrees, and the volume resistivity was 1×10° Ω-cm. The elvel of transfer current required for transfering the image from the photoconductor drum 1 to the intermediate transfer body 5 is about 5 μA, and this level of current was obtained by adding a voltage of +500 V to the cored bar 5b.
- [0179] The four-color toner color combination latent image formed on the intermediate transfer body 5 is transferred to an object transferring material such as a paper by a transfer roller?, and is thereafter fixed by a heal-fixation appearable. H. The transfer roller is provided thereon the core metal To with the outside diameter of 10 mm on which an exist layer 7a is formed by coating of a foam of eithylene-propylene-diene based tridimensional copolymer (EPDM) dispersing carbon sufficiently therein as a conductivity producing material. The layer had a volume specific resistance of 1 x 10°S, cm and a hardness degree of 35° as measured in accordance with "JIS K-6301". In addition, a voltage was
 - applied to this transfer roller 7 to pass a transfer current of 15 µA thereIntrough.

 [0180] In the apparatus shown in FIG. 2, a fixation apparatus of heated roll type having no oil coating mechanism shown in FIGS. 8 and 7 was used in the heat-fixation apparatus H. The both upper and lower rollers of the fixation apparatus used here had surface layers made of fluorino based resin. In addition, the diameter of the roller was 50 mm. The fixation temperature for fixation was 160°C, and the highping width was set at 7 mm. Furthermore, a transfer residual toner on the photoconductor drum 1, which was collected by cleaning, was transported to a developing device by a reuse mechanism for reuse.

Evaluation

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[0181] Two-component type developers produced using the toners of Examples 5 to 22 and two-component type developers produced using toners of Comparative Examples 1 to 6 were used, respectively, to perform printuit testing at a printout trate of 8 sheets (A4 size) per minute while the developer was supplied one after another in a monochromatic intermittent mode (namely a mode in which the developing device is stopped for 10 seconds for each printout to accelerate the degradation of a toner in a preliminary operation during restart of the device) at a normal temperature and normal humidity (25°C, 60% Rh¹) and a high temperature and high humidity (30°C, 80% Rh¹) under the conditions described above, and resulting printout images were evaluated for the following items. The evaluation results are shown together in Table 12.

50 Evaluation of printout images

1. Image density

[0182] Images were printed out on a predetermined number of normal copying papers (75 g/m²), and the image density was evaluated according to the level at which the density of the image from the final printout was retained with respect to the density of the initial image. Furthermore, for the measurement of image density, a Macbeth reflective densitometer (manufactured by Macbeth Co., Ltd.) was used to measure a density relative to that of the printout image on a white cround with the density of orionial coop or equal to 0.00.

EP 1 336 635 A1

- A: Excellent (image density from the final printout is 1.40 or greater)
- B: Good (image density from the final printout is 1.35 or greater and lower than 1.40)
- C: Usable (image density from the final printout is 1.00 or greater and lower than 1.35)
- D: Unusable (image density from the final printout is lower than 1.00)

Image fog

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[D183] Images were printed out on a predetermined number of normal copying papers (76 g/m²), and the image frog was evaluated with a solid white lemage from the final printout. Sepecifically, the oveluation was made as foliow: the worst value of the reflective density of the white ground after printing and the average reflective density of the paper before printing, as measured using a reflective densitometer (Palc LT-CaBS manufactured by Tokyo Denshoku Co., Ltd.), were defined as Ds and Dr, respectively, (Ds-Dr) was calculated from these values as a fog level to make an evaluation according to the following criterion.

- A: Excellent (fog level is 0% or higher and lower than 1.5%)
 - B: Good (fog level is 1.5% or higher and lower than 3.0%)
 - C: Usable (fog level is 3.0% or higher and lower than 5.0%)
 - D: Unusable (fog levei is 5.0% or higher)

20 3. Transferability

[0184] Solid black Images were printed out on a predetermined number of normal copying papers (75 g/m²), and the image dislocation level of the image from the final printout was visually observed to make an evaluation according to the following oriterion.

- A: Excellent (almost not observed)
- B: Good (slightly observed)
- C: Usable
- D: Unusable

[0185] In addition, in Examples 23 to 40 and Comparative Examples 7 to 12, occurrences of scares and sticking residual toners on the surfaces of the photoconductor drum and intermediate transfer body, and their influence on printout images (matching with the image forming apparatus) were usually evaluated after 5000 images were outputted, and as a result, scars and sticking residual toners on the surfaces of the photoconductor drum and intermediate transfer body were not observed, and thus matching with the image forming apparatus was excellent for the system using two-component type developers of Examples 23 to 40. For the system using two-component type developers of Comparative Examples 27 to 12, on the other hand, sticking toners were observed on the surface of the intermediate transfer body, and there was a problem in matching with image formation apparatus such that longitudinally striped defects occurred on the image.

Toble 1

				Ta	able 12				
45	Examples	Two component type developer	Norma	l temperature humidit	e and normal y	High temperature and high humidity			
			Image density	Image fog	Transferability	image density	Image fog	Transferability	
50	23	Blue 1	Α	Α	Α	Α	AA	Α	
	24	Blue 2	Α	Α	Α	Α	Α	Α	
	25	Blue 3	В	Α	Α	В	В	Α	
55	26	Yellow 1	Α	Α	Α	Α	Α	Α	
	27	Yellow 2	Α	Α	Α	Α	Α	Α	
	28	Yellow 3	В	A	Α	В	В	Α	

EP 1 336 635 A1

Table 12 (continued)

Exe	amples	Two component type developer	Norma	l temperatur humidit	e and normal y	High temperature and high humidity			
			Image density	Image fog	Transferability	Image density	Image fog	Transferability	
29		Black 1	Α	Α	A	Α	Α	Α	
30		Black 2	Α	Α	Α	А	Α	Α	
31		Black 3	Α	В	В	В	В	В	
32		Red 1	Α	Α	Α	Α	Α	Α	
5 33		Red 2	Α	Α	Α	А	Α	Α	
34		Red 3	Α	Α	В	В	В	В	
35		Black 5	Α	Α	A	Α	Α	Α	
36		Black 6	Α	Α	Α	Α	Α	Α	
37		Black 7	В	Α	Α	В	В	Α	
38		Black 9	Α	Α	Α	Α	Α	A	
39		Black 10	Α	Α	Α	Α	Α	Α	
5 40		Black 11	В	В	В	В	В	В	
	parative ple 7	Blue 4	D	D	D	D	D	D	
	parative	Yellow 4	D	D	D	D	D	D	
	parative	Black 4	С	С	D	С	D	D	
	parative npie 10	Red 4	С	С	D	С	D	D	
	parative	Black 8	С	С	D	D	D	D	
	parative	Black 12.	С	С	D	С	D	D	

Examples 41 to 43 and Comparative Examples 13 to 15

[0188] For carrying out the image formation methods of Examples 41 to 43 and Comparative Examples 13 to 15.
The toners obtained in Examples 5, 8 and 11 and Comparative examples 1 to 3 were used, respectively, as devolerely, and as devolerely, as devolerely, as devolerely, as devolerely, and devolerely, and devolerely, and devolerely, as devolerely, and devolerely, as devolerely, and devol

[0187] In the image forming apparatus shown in FIG. 4, the surface of the photoconductor drum 20 is electrically charged by a primary charge roller 27. A rubber roller (diameter 12 mm, abutment pressure 50 g/cm) coated with a nylon resin and having conductive carbon dispersed therein was used for the primary charge roller 27, and an electrostatic latent image with a dark area potential VD of -700 V and a light area potential VL of -200 V was formed on the electrostatic latent image carrier (photoconductor drum 20) by Islaser exposure (600 dp. not shown). As a toner

FP 1 336 635 A1

carrier, a development sleeve 28 having a roughness degree Ra of 1.1 with the surface coated with a resin having a carbon black dispersed therein was used.

[0188] An enlarged sectional view of the principal part of the development apparatus for one-component type developers used in Examples 41 to 43 and Comparative Examples 13 to 15 is shown in FIG. 5. For conditions for developing electrostatic latent images, the speed of the development sleeve 28 was set at a speed 1.1 times as high as the movement speed of the surface of the photoconductor drum 20 opposite thereto, and the space α between the photoconductor drum 20 and the development sleeve 28 (between 5 and D) was 270 µm. For the member for controlling the thickness of the toner, an abutting urethane rubber blade 29 was used. In addition, the set temperature of the heat-fixation apparatus for fixing a toner image was 160°C. Furthermore, for the fixation apparatus, a fixation apparatus α shown in FIG. 5.8 and 7 was used.

Shown in PGS. a did 7 was used.

(D189] As described above, under the condition of normal temperature and normal humidity (25°C, 60%RH), images were printed out on up to 30,000 sheets at a printout rate of 8 sheets (A4 size) per minute while the toner was supplied one after another in a continuous mode (namely, a mode in which the development device is not stopped, and thereby consumption of the toner is promoted), and the densities of resulting printout images were measured to evaluate the durability of the image according to the following criterion. In addition, the image from the 10,000 th printout was observed to make an evaluation about image forg according to the following criterion. At the same time, situations of the components constituting the image forming apparatus after the durability testing were observed to evaluate matching between each component and the above described toner. The results thereof are shown toestlerin in Table 13.

20 Change in image density during endurance

[0190] Images were printed out on a predetermined number of normal copying papers (75 g/m²), and the image density was evaluated according to the level at which the density of the image from the final printout was retained with respect to the density of the initial image. Furthermore, for the measurement of image density, a Macbeth reflective densitometer (manufactured by Macbeth Co., Ltd.) was used to measure a density relative to that of the printout image on a white ground with the density of original copy equal to 0.0.

- A: Excellent (Image density from the final printout is 1.40 or greater)
- B: Good (image density from the final printout is 1.35 or greater and lower than 1.40)
- C: Usable (image density from the final printout is 1.00 or greater and lower than 1.35)
- D: Unusable (image density from the final printout is lower than 1.00)

Image fog

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[0191] Images were printed out on a predetermined number of normal copying papers (75 g/m²), and the image log was evaluated with a solid white image from the final printout. Specifically, the evaluation was made as follow: the worst value of the reflective density of the white ground after printing and the average reflective density of the paper before printing, as measured using a reflective densitometer (Reflectometer ODEL TC-6DS manufactured by Tokyo Denshoku Co., Ltd.), were defined as Ds and Dr, respectively, (Ds-Dr) was calculated from these values as a fog level to make an evaluation according to the following criterion.

- A: Excellent (fog level is 0% or higher and lower than 1.5%)
- B: Good (fog level is 1.5% or higher and lower than 3.0%)
- C: Usable (fog level is 3.0% or higher and lower than 5.0%)
 - D: Unusable (fog level is 5.0% or higher)

Evaluation of matching with image forming apparatus

1. Matching with development sleeve

[0192] After the printout testing was completed, the situation of residual toners sticking to the surface of the development sleeve and their influence on the printout image were visually evaluated.

- A: Excellent (not observed)
- B: Good (almost not observed)
- C: Usable (sticking residual toners are observed but the influence on the image is not significant)
- D: Unusable (sticking of residual toners is significant, causing unevenness in the image)

FP 1 336 635 Δ1

- 2. Matching with photoconductor drum
- [0193] Occurrences of scars and sticking residual toners on the surface of the photoconductor drum and their influence on the printout image were evaluated visually.
- A: Excellent (not observed)

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- B: Good (slightly observed but no influence on the image)
- C: Usable (sticking residual toners and scars are observed but the influence on the image is not significant)
- D: Unusable (sticking of residual toners is significant, causing longitudinal striped defects in the image)
- 3. Matching with fixation apparatus
- [0194] The surface situation of the fixation film was observed, and the results of surface characteristics and occurrences of sticking residual toners were collectively averaged to evaluate the durability of the film.
 - (1) Surface characteristics
 - [0195] Occurrences of scares and flaking on the fixation film were visually observed and evaluated after the printout testing was completed.
 - A: Excellent (not observed)
 - B: Good (almost not observed)
 - C: Usable
- D: Unusable
 - (2) Situation of sticking toners
 - [0196] The situation of residual toners sticking to the surface of the fixation film was visually observed and evaluated after the printout testing was completed.
 - A: Excellent (not observed)
 - B: Good (almost not observed)
 - C: Usable D: Unusable

5			tus	Toner fixation	A	¥	ď	۵	۵	۵
10		apparatus	Fixation apparatus	surface	A	¥	V	Q	Q	Q
15		Evaluation of matching with other apparatus	Photoconduct or drum		A	A	٧	٥	۵	۵
20		Evaluation of me	Development sleeve		V	A	٧	۵	۵	۵
25	Table 13		10 thousands fogged images		4	4	4	۵	۵	۵
30 35	Tab		ance	30 thousands	A	A	¥	۵	۵	۵
40		t image	Change in image density during endurance	Thousand 10 thousands 30 thousands	4	4	∢	۵	۵	۵
45		Evaluation of printout image	e in image der		ď	4	Ą	۵	۵	υ
		Evalua	Chang	Initial	<	4	∢	O	O	8
50		Toner			Blue 1	Yellow 1	Black 1	Blue 4	Yellow 4	Black 4
55		Examples			41	42	43	Comparative Example 13	Comparative Example 14	Comparative Example 15

Example 44

[0197] Printout testing was performed while the blue toner (1) of Example 5 was supplied one after another in a continuous mode (namely, a mode in which the development device is not stopped, and thereby consumption of the toner is promoted) in the same manner as Example 41 except that the toner reuse mechanism of the image forming apparatus of FIG. 4 was removed and that the printout rate was set at the level of 16 sheets (Avi size) per minute. The resulting printout images and the matching with the image evaluating apparatus used were evaluated for the same items as Examples 41 to 43 and Comparative Examples 13 to 15. As a result, satisfactory results were obtained for all the items.

Claims

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1. A polyhydroxyalkanoate comprising a unit of formula (1):

wherein R is selected from the group consisting of OH, a hatogen atom, ONa, OK, OCH₃ and OC₂H₃. A represents a substituted or unsubstituted alphatic hydrocarbon structure; m is an integer number selected from 1 os and in the case where a plurality of units exist in the same molecule, R, A and m in one unit can be different from them in another unit respectively.

2. The polyhydroxyalkanoate according to claim 1, comprising a unit of formula (2):

wherein R_i is H or CH_3 , R_i is selected from the group consisting of OH, a helogen atom, ONa, OK, OCH_3 and OCH_3 , R_i and CC_3 each represent a substituted or unsubstituted alphalet hydrocarbon structure, in it is an integer number selected from 1 to R_i and in the case where a plurality of units exist in the same molecule, R_1 , R_2 , R_1 , C_1 and m in one unit can be different from them in another unit respectively.

3. The polyhydroxyalkanoate according to claim 2, comprising a unit of formula (3):

FP 1 336 635 Δ1

wherein R₃ is H or CH₃; R₄ is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and OC₂H₅; B₂ and C₂ each represent a straight-chain or branched alkylene group having 1 to 8 carbon atoms; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R₃, R₄, B₂, C₂ and m in one unit can be different from them in another unit respectively.

4. The polyhydroxyalkanoate according to claim 3, comprising a unit of formula (4):

wherein R_3 is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH $_3$ and OC $_2$ H $_5$; m is an integer number selected from 1 to 8, and in the case where a plurality of units exist in the same molecule, R_3 and m in one unit can be different from them in another unit respectively.

- The polyhydroxyalkanoate according to claim 1, wherein m in formula (1) is an integer selected from the group consisting of 4, 6 and 8.
- 6. The polyhydroxyalkanoate according to claim 1, wherein m in formula (1) is an integer of 3 or 5.
- The polyhydroxyalkanoate according to claim 1, comprising at least one of a 3-hydroxy-ω-alkanoic acid unit of formula (5):

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wherein is an integer number selected from 1 to 8; R_0 contains a residue having a phenyl structure or at theirly structure; and in the case where a plurality of units exist in the same molecule, n and R_0 in one unit can be different from them in another unit respectively and

a 3-hydroxy-ω-cyclohexylalkanoic acid unit of formula (6):

wherein R_7 is a substituent group in the cyclohexyl group selected from the group consisting of H, CN, NO₂, a halogen atom, CH₃, C₂H₃, C₃H₇, CF₃, C₂F₅ and C₃F₇; k is an integer number selected from 0 to 8; and in the case where a plurality of units exist in the same molecule, k and R₇ in one unit can be different from them in another unit respectively.

The polyhydroxyalkanoate according to claim 7, wherein R₆ in formula (5) is selected from the group consisting
of the groups of the following formulas (7), (8), (9), (10), (11), (12), (13), (14), (15), (16) and (17):

unsubstituted or substituted phenyl groups of formula (7):

wherein R_{θ_0} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO $_2$, CH $_3$, C2H $_3$, CQC ON $_{\theta_0}$ (R_{θ_0} represents any one selected from the group consisting of H, Na and K), CF $_3$, C2F $_3$ and C3F $_3$, and in the case where a plurality of units exist in the same molecule, R_{θ_0} in one unit can be different from that in another unit; unsaturated or saturated otheroxy croups of formula (8):

wherein R_S represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, CH₅, Cg+1₅, Cg+1₅, CG+3, Cg-5, Cg-5, and Cg-7, and in the case where a plurality of units exist in the same molecule, R_S in one unit can be different from that in another unit; unsubstituted or substituted benzoyl groups each of formula (9):

wherein R_{10} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, CH₃, C₂H₃, C₃H₃, C₃F₃, C₃F₃ and C₃F₇, and in the case where a plurality of units exist in the same molecule, R_{10} in one unit can be different from that in another unit; unsubstituted or substituted phenylsulfanyl groups of formula (10):

wherein R_{Ha} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, ON, NO_2 , $COOR_{Hb}$, OP_{2H_0} (R_{Hb} represents any one selected from the group consisting of H, at N, A, C, H and C_{2H}, and R₁ represents any one selected from the group consisting of OH, ON, ON, a halogen atom, OCH_3 and OC_2H_3 , OP_3

wherein R_{12a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, COOR_{12b}, SO₂R_{12c} (R_{12b} represents any one selected from the group consisting of H, A, N, K, CH₃ and C₂H₅, and R_{12c} represents any one selected from the group consisting of OH, ONa, OK, a halogen atom, OCH₃ and OC₂H₅), CH₃, C₂H₅, C₃H₇, (CH₃)₂·CH and (CH₃)₃·C, and in the case where a plurality of units exist in the same molecule, R_{12a} in one unit can be different from that in another unit; 2-thienvi croup of formula (12):

2-thienvisulfanyl group of formula (13):

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2-thienylcarbonyl group of formula (14):

unsubstituted or substituted phenylsulfinyl groups of formula (15):

wherein R_{13a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, COOR₁₃₀(R₁₃₀ represents any one selected from the group consisting of H, A, K, CH₃ and C₃H₃, SO₂R₁₃₀(R₁₃₀, represents any one selected from the group consisting of OH, ONa, OK, a halogen atom, OCH₃ and OC2₂H₃), CH₃, C₂H₃, C₃H₃, (CH₃)₂-CH and (CH₃)₂-C, and in the case where a plurality of units exist in the same molecule, R_{13a} in one unit can be different from that in another unit; unsubstituted or substituted phenylsufforly (groups of formula (16):

wherein R_{14a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, COOR₁₄₆(R₁₄₆, represents any one selected from the group consisting of H, Na, K, CH₃ and C_PH₃, SO₂P₁₄₆(R₁₄₆, represents any one selected from the group consisting of CH, Na, OK, a halogen atom, OCH₃ and OC₂H₃), CH₃, C₂H₃, C₃H₇, (CH₃)₂·CH and (CH₃)₂·C, and in the case where a plurality of units exist in the same molecule, R_{14a} in one unit can be differend from that in another unit; and (phenyimethylyoxy groups of formula (17).

and in the case where a plurality of units exist in the same molecule, R₆ in one unit of formula (5) can be different from that in another unit.

9. The polyhydroxyalkanoate according to claim 1, wherein the number-average molecular weight is in the range of

from 1000 to 1000000.

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10. A method of producing polyhydroxyalkanoate comprising a unit of formula (1):

wherein R is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and OC₂H₅: A represents a substituted or unsubstituted aliphatic hydrocarbon structure; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R, A and m in one unit can be different from them in another unit respectively. which comprises the step of:

reacting a polyhydroxyalkanoate comprising a unit of formula (18):

wherein m is an integer number selected from 1 to 8, and in the case where a plurality of units exist in the same molecule, m in one unit can be different from that in another unit, with at least one type of compounds of formula (19):

wherein R_{15} is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH $_3$ and OC $_2$ H $_5$ and A_1 is a substituted or unsubstituted aliphatic hydrocarbon structure, and in the case where a plurally of types of compounds exist in the same molecule, R_{15} and A_1 in one unit can be different from them in another unit respectively.

11. A method of producing polyhydroxyalkanoate comprising a unit of formula (2):

When the properties of the pr

reacting a polyhydroxyalkanoate comprising a unit of formula (18):

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wherein m is an integer number selected from 1 to 8, and in the case where a plurality of units exist in the same molecule, m in one unit can be different from that in another unit, with at least one type of compounds of formula (20):

$$HS-B_3 - N-C_3-SO_2R_{17}$$
(20)

wherein R_{16} is H or CH_3 ; R_{17} is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and CO_2H_5 ; B_3 and CO_3 are selected from substituted or unsubstituted aliphatic hydrocarbon structures; and in the case where a plurality of types of compounds exist in the same molecule, R_{16} , R_{17} , R_3 and C_3 in one unit can be different from them in another unit respectively.

- 12. The method according to claim 10, wherein the reacting step is carried out under basic condition.
- 13. The method according to claim 12, wherein at least one selected from the group consisting of dimetylamine, diethylamine, trimethylamine, triethylamine, morpholine, piperidine, sodium hydroxide, potassium hydroxide, sodium carbonate, potassium carbonate, sodium methylate and sodium ethylate is used as a basic catalyst in the reacting step.
- 14. A charge controlling agent controlling the charged state of powder, the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (1):

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wherein R is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and OC₂H₅: A represents a substituted or unsubstituted aliphatic hydrocarbon structure; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R, A and m in one unit can be different from them in another unit respectively.

15. The charge controlling agent controlling the charged state of powder according to claim 14, the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (2):

wherein R_1 is H or CH_3 : R_2 is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH_3 and OC_2H_3 , B_1 and C_1 each represent a substituted or unsubstituted aliphatic hydrocarbon structure; m is an integer number selected from 1 to B_1 and in the case where a plurality of units exist in the same molecule, R_1 , R_2 , B_1 , C_1 and m in one unit can be different from them in another unit respectively.

16. The charge controlling agent controlling the charged state of powder according to claim 15, the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (3):

wherein R_3 is H or CH_3 : R_4 is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and C_2 -plant represent a straight-chain or branched alkylene group having 1 to 8 carbon atoms; m is an integer number selected from 1 to 8; and in the case where a plurality of units exist in the same molecule, R_3 . R_4 , B_5 , C_5 , and m in one unit can be different from them in another unit respectively.

17. The charge controlling agent controlling the charged state of powder according to claim 16, the charge controlling agent containing polyhydroxyalkanoate comprising a unit of formula (4):

wherein R₅ is selected from the group consisting of OH, a halogen atom, ONa, OK, OCH₃ and OC₂H₅; m is an integer number selected from 1 to 8, and in the case where a plurality of units exist in the same molecule, R₅ and m in one unit can be different from them in another unit respectively.

18. The charge controlling agent according to claim 14, comprising at least one of a 3-hydroxy-ω-alkanoic acid unit of formula (5):

wherein n is an integer number selected from 1 to 8; Rs contains a residue having a phenyl structure or a thienyl structure; and in the case where a plurality of units exist in the same molecule, n and Re in one unit can be different from them in another unit respectively

a 3-hydroxy-w-cyclohexylalkanoic acid unit of formula (6):

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wherein R7 is a substituent group in the cyclohexyl group selected from the group consisting of H, CN, NO2, a halogen atom, CH3, C2H5, C3H7, CF3, C2F5 and C3F7; k is an integer number selected from 0 to 8; and in the case where a plurality of units exist in the same molecule, k and R, in one unit can be different from them in another unit respectively.

19. The charge controlling agent according to claim 18, wherein R₆ in chemical formula (5) is selected from the group consisting of the groups of the following formulas (7), (8), (9), (10), (11), (12), (13), (14), (15), (16) and (17):

unsubstituted or substituted phenyl groups of formula (7):

wherein Rg, represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO2, CH3, C2H5, C3H7, CH=CH2, COORgh (Rgh represents any one selected from the group consisting of H, Na and K), CF₃, C₂F₅ and C₃F₇, and in the case where a plurality of units exist in the same molecule, R_{8a} in one unit can be different from that in another unit;

unsaturated or saturated phenoxy groups of formula (8):

wherein R₃ represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, CH₃, CH₃, C₃H₃, SCH₃, CF₅, and C₃F₇, and in the case where a plurality of units exist in the same nolecule, R₃ in one unit can be different from that in another unit; unsubstituted or substituted because of formula (9):

wherein R_{ij} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO_2 , CH_3 , C_2H_2 , C_3H_2 , C_3H_2 , C_3H_2 , and in the sase where a plurality of units exist in the same molecule, R_{ij} in one unit can be different from that in another unit; unsubstituted or substituted hereinvalled and consists of simple of the same consists of the same cons

wherein R_{11a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, No $_2$, COOR_{11b}, So $_2$ R $_{11c}$ (R $_{11b}$ represents any one selected from the group consisting of OH, ONa, OK, a halogen atom, OCH $_3$ and CQ $_{11b}$, and III, represents any one selected from the group consisting of OH, ONa, OK, a halogen atom, OCH $_3$ and OC $_2$ H $_3$), CH $_3$, C $_2$ H $_5$, C $_2$ H $_5$, C $_3$ H $_7$, (CH $_3$) $_2$ CH and (CH $_3$) $_3$ C, and in the case where a plurality of units exist in the same molecule, R $_{11a}$ in one unit can be different from that in another unit; unsubstituted or substituted (oherwinethyt) sulfaying droups of formula (11):

wherein R_{12a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, COOR_{12b}, SO₂R_{12c} (R_{12b} represents any one selected from the group consisting of OH, Na, K, CH₃ and C₂H₅, and R_{12c} represents any one selected from the group consisting of OH, ONa, OK, a halogen atom, OCH₃ and OC₂H₅), CH₃, C₂H₅, C₃H₇, (CH₃)₂-CH and (CH₃)₃-C, and in the case where a plurality of units exist in the same molecule, R_{12a} in one unit can be different from that in another unit; 2-thineny (group of formula (12):

EP 1 336 635 A1

2-thienylsulfanyl group of formula (13):

2-thienylcarbonyl group of formula (14):

unsubstituted or substituted phenylsulfinyl groups of formula (15):

wherein R_{13a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO_2 , $COOR_{136}(R_{136}$, represents any one selected from the group consisting of H, Na, K, CR_1 , and C_2 H₃, SO_2 R₁₃₆(R₁₃₆, represents any one selected from the group consisting of OH, ON_3 , OK, a halogen atom, OCH_3 and OC_2 H₃), CH_3 , C_2 H₃, C_3 H₇, $(CH_3)_2$ -CH and $(CH_3)_3$ -C, and in the case where a plurality of units exist in the same molecule, R_{13a} in one unit can be different from that in another unit; unsubstituted or substituted phenylsuilforly groups of formula (15).

wherein R_{14a} represents a substituent group in the aromatic ring selected from the group consisting of H, a halogen atom, CN, NO₂, COOR₁₄₈(R₁₄₈, represents any one selected from the group consisting of H, AC, H3 and C₂H3), SO₂R₁₄₆(R₁₄₆, represents any one selected from the group consisting of OH, ONa, OK, a halogen atom, OCH₃ and OC₂H3), CH₃-C₂H3, C₃H7, (CH₃)₂-CH and (CH₃)₃-C, and in the case where a plurality of units exist in the same molecule, R_{14a} in one unit can be different from that in another unit; and (pheny/methyl)oxy groups of formula (17):

EP 1 336 635 A1

, and in the case where a plurality of units exist in the same molecule, R_6 in one unit can be different from that in another unit.

- The charge controlling agent according to claim 14, wherein said powder is an electrostatic latent image developing toner.
- The charge controlling agent according to claim 14, wherein the number-average molecular weight of said polyhydroxvalkanoate is in the range of from 1000 to 1000000.
- 22. A toner binder for use in an electrostatic latent image developing toner, the toner binder containing the charge controlling agent according to claim 14.
- An electrostatic latent image developing toner containing a binder resin, a coloring agent and the charge controlling
 agent according to claim 14.
 - 24. A method for forming an image which comprises the steps of:
 - applying a voltage to an electrification member from the outside to electrify an electrostatic latent image carrier.
 - forming an electrostatic latent image on the electrified electrostatic latent image carrier,
 - developing the electrostatic latent image by an electrostatic latent image developing toner to form a toner image on the electrostatic latent image carrier, transferring the toner image on the electrostatic latent image carrier to a record material, and
 - fixing the toner image on the record material by heat, wherein an electrostatic latent image developing toner according to claim 23 is used.
- 25. The method according to claim 24, wherein said transferring step comprises a first transferring step of transferring step of transferring step of transferring step of transferring the toner image on the electrostatic latent image carrier to an intermediate transfer body to the record material.
 - 26. An image forming apparatus comprising a means for applying a voltage to an electrification member from the outside to electrify an electrostatic latent image carrier, a means for forming an electrostatic latent image on the electrified electrostatic latent image to make the electrostatic latent image developing toner to form a toner image on the electrostatic latent image carrier, a means for transferring the toner image on the electrostatic latent image carrier, a means for transferring the toner image on the electrostatic latent image carrier to a record material, and a means for fixing the toner image on the record material by heat, wherein an electrostatic latent image developing toner according to claim 23 is used.
 - 27. The image forming apparatus according to claim 26, wherein said transferring means comprises a first transferring means for transferring the electrostatic latent image carrier to an intermediate transfer body and a second transferring means for transferring the toner image on the intermediate transfer body to the record material.

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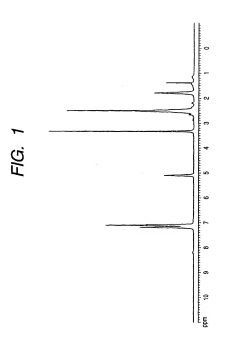


FIG. 2

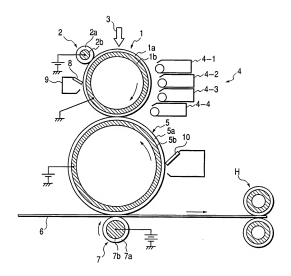


FIG. 3

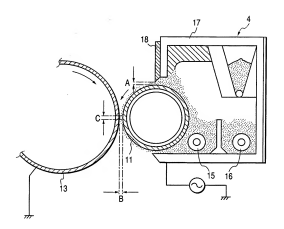


FIG. 4

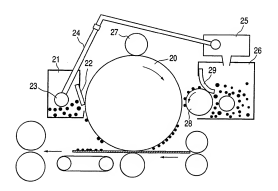
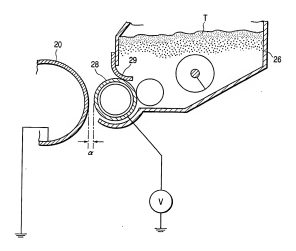


FIG. 5



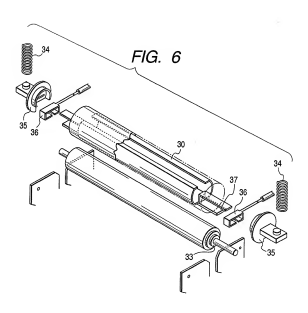


FIG. 7

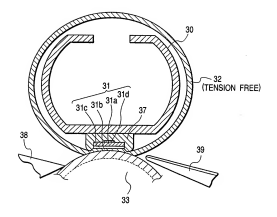
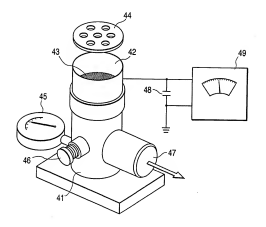


FIG. 8





European Patent

EUROPEAN SEARCH REPORT

Application Number EP 03 00 3419

	DOCUMENTS CONSID	ERED TO BE RELEVANT			
Category	Citation of document with I	ndication, where appropriate, seges	Relevant to claim	CLASSIFICATION OF THE APPLICATION (IntCL7)	
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		*		TECHNICAL FIELDS SEARCHED (Int.Cl.7)	
				C086 C12P G036	
	The present search report has	· ·			
	Place of search	Date of completion of the search		Extensor	
	THE HAGUE	26 May 2003		ocker, L	
X : parti Y : parti docu A : tech O : non	ATEGORY OF CITED DOCUMENTS cutarly relevant if taken alone cutarly relevant if combined with anot meet of the same category nological background -written disclosure mediate document		the application rother reasons		

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 03 00 3419

This annex lists the patent samily membersrelating to the patent documents clied in the above—mentioned European search report. The members are as contained in the European Patent Office ECP file on The European Patent Office is in no way liabile to these particulars which are merely given for the purpose of information.

26-05-2003

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For more details about this annex; see Official Journal of the European Patent Office, No. 12/82